

WORLD IN TELLECTUAL PROPERTY ORGANIZATION



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(54) Title: FLK-1 IS	A REC	EPTOR FOR	VASCUL	ar ei	VD(OTHELIAL GROWTH FACTOR		
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TKR-C								
FLK-1 KDR TKR-C	926	GKDYVGELSVDLKRRLDSITSSQSSASSGFVEEKSLSDVEEEEASEELYKDFLTLEHLIC ———AIP—————————————————————————————————						
FLK-1 KDR TKR-C	986	YSFQVAKCMEFLASRKCIHRDLAARNILLSEKNVVKICDFGLARDIYKDPDYVRKGDARL						
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(57) Abstract

The present invention relates to the use of ligands for the Flk-1 receptor for the modulation of angiogenesis and vasculogenesis. The invention is based, in part, on the demonstration that Flk-1 tyrosine kinase receptor expression is associated with endothelial cells and the identification of vascular endothelial growth factor (VEGF) as the high affinity ligand of Flk-1. These results indicate a major role for Flk-1 in the signaling system during vasculogenesis and angiogenesis. Engineering of host cells that express Flk-1 and the uses of expressed Flk-1 to evaluate and screen for drugs and analogs of VEGF involved in Flk-1 modulation by either agonist or antagonist activities is described. The invention also relates to the use of FLK-1 ligands, including VEGF agonists and antagonists, in the treatment of disorders, including cancer, by modulating vasculogenesis and angiogenesis.

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Flk-1 IS A RECEPTOR FOR VASCULAR ENDOTHELIAL GROWTH FACTOR

1. INTRODUCTION

The present invention relates to the use of ligands for the FLK-1 receptor for the modulation of angiogenesis and vasculogenesis. The invention is based, in part, on the demonstration that Flk-1 tyrosine kinase receptor expression is associated with endothelial cells and the identification of vascular endothelial growth factor (VEGF) as the high affinity ligand of Flk-1. These results indicate a major role for Flk-1 in the signaling system during vasculogenesis and angiogenesis. Engineering of host cells that express Flk-1 and the uses of expressed Flk-1 to evaluate and screen for drugs and analogs of VEGF involved in Flk-1 modulation by either 15 agonist or antagonist activities is described.

The invention also relates to the use of FLK-1 ligands, including VEGF agonists and antagonists, in the treatment of disorders, including cancer, by modulating vasculogenesis and angiogenesis.

2. BACKGROUND OF THE INVENTION

Receptor tyrosine kinases comprise a large family of transmembrane receptors for polypeptide growth factors with diverse biological activities. Their intrinsic 25 tyrosine kinase function is activated upon ligand binding, which results in phosphorylation of the receptor and multiple cellular substrates, and subsequently in a variety of cellular responses (Ullrich A. and 30 Schlessinger, J., 1990, Cell 61:203-212).

A receptor tyrosine kinase cDNA, designated fetal liver kinase 1 (Flk-1), was cloned from mouse cell populations enriched for hematopoietic stem and progenitor cells. The receptor was suggested to be involved in hematopoietic stem cell renewal (Matthews

for growth.

et al., 1991, Proc. Natl. Acad. Sci. USA 88:9026-9030).
Sequence analysis of the Flk-1 clone revealed
considerable homology with the c-Kit subfamily of
receptor kinases and in particular to the Flt gene
product. These receptors all have in common an
extracellular domain containing immunoglobulin-like
structures.

The formation and spreading of blood vessels, or vasculogenesis and angiogenesis, respectively, play important roles in a variety of physiological processes such as embryonic development, wound healing, organ regeneration and female reproductive processes such as follicle development in the corpus luteum during ovulation and placental growth after pregnancy.

15 Uncontrolled angiogenesis can be pathological such as in the growth of solid tumors that rely on vascularization

Angiogenesis involves the proliferation, migration and infiltration of vascular endothelial cells, and is likely to be regulated by polypeptide growth factors. Several polypeptides with in vitro endothelial cell growth promoting activity have been identified. Examples include acidic and basic fibroblastic growth factor, vascular endothelial growth factor and placental growth factor. Although four distinct receptors for the different members of the FGF family have been characterized, none of these have as yet been reported to be expressed in blood vessels in vivo.

While the FGFs appear to be mitogens for a large
number of different cell types, VEGF has recently been
reported to be an endothelial cell specific mitogen
(Ferrara, N. and Henzel, W.J., 1989, Biochem. Biophys.
Res. Comm. 161:851-858). Recently, the fms-like tyrosine
receptor, flt, was shown to have affinity for VEGF

35 (DeVries, C. et al., 1992, Science 255:989-991).

3. SUMMARY OF THE INVENTION

The present invention relates to the use of ligands for the FLK-1 receptor for the modulation of angiogenesis and vasculogenesis. The present invention is based, in part, on the discovery that the Flk-1 tyrosine kinase receptor is expressed on the surface of endothelial cells and the identification of vascular endothelial growth factor (VEGF) as the high affinity ligand of Flk-1. The role of endothelial cell proliferation and migration during angiogenesis and vasculogenesis indicate an important role for Flk-1 in these processes. The invention is described by way of example for the murine Flk-1, however, the principles may be applied to other species including humans.

Pharmaceutical reagents designed to inhibit the Flk-1/VEGF interaction may be useful in inhibition of tumor growth. VEGF and/or VEGF agonists may be used to promote wound healing. The invention relates to expression systems designed to produce Flk-1 protein and/or cell lines which express the Flk-1 receptor. Expression of soluble recombinant Flk-1 protein may be used to screen peptide libraries for molecules that inhibit the Flk-1/VEGF interaction. Engineered cell lines expressing Flk-1 on their surface may be advantageously used to screen and identify VEGF agonists and antagonists.

4. BRIEF DESCRIPTION OF THE FIGURES

FIG. 1. Comparison of the Flk-1 amino acid
sequence with related RTKs. Amino acid sequence
comparison of Flk-1 with human KDR and rat TKr-C. A
section of the sequence which is known for all three
receptors is compared and only differences to the Flk-1
sequence are shown.

- FIG. 2. Northern blot analysis of Flk-1 gene expression. (A) Expression of Flk-1 RNA in day 9.5 to day 18.5 mouse embryos. Samples (10 μg) of total RNA from whole mouse embryos were analyzed in each lane.
 5 Positions of 28S and 18S ribosomal RNAs are marked. (B) Expression of Flk-1 mRNA in postnatal day 4 and adult brain in comparison with capillary fragments from postnatal day 4 brain. 1μg of poly (A⁺) RNA was loaded on each lane. The 5' 2619 bp of the Flk-1 cDNA were used as a probe. Control hybridization with a GAPDH cDNA probe is shown in the lower panel.
- FIG. 3. Abundant Flk-1 gene expression in embryonic tissues. In situ hybridization analysis of Flk-1 expression in day 14.5 mouse embryo. (A) Bright field illumination of a parasagittal section through the whole embryo hybridized with a 35S-labeled antisense probe (5' 2619 bp). (B) Dark field illumination of the same section. (C) Control hybridization of an adjacent section with a sense probe. Abbreviations: Ao, aorta; At, atrium; L, lung; Li, liver; Ma, mandible; Mn, meninges; Ms. mesencephalon; T, telencephalon; V, ventricle; Vt, vertebrae.
- FIG. 4. Expression of Flk-1 RNA in embryonic organs is restricted to specific cells. Expression of Flk-1 RNA in a day 14.5 mouse embryo at higher magnification. (A) The heart region was probed with a "S-labeled antisense probe. (B) Adjacent section hybridized with the sense probe. (C) Part of the aorta wall shown on the cellular level. The endothelial cell-layer is indicated by an arrow. (D) The lung, probed with the Flk-1 antisense probe. (E) Control hybridization of an adjacent section hybridized with the sense probe. Abbreviations: At, atrium; B, bronchus; Ed, endothelial cell layer; En, endocardium; L, lung, Li,

liver; Lu, lumina of the aorta; Ml, muscular; My, myocardium.

- FIG. 5. Flk-1 gene expression in the brain of the developing mouse. In situ hybridization analysis of Flk-5 1 gene expression in the brain at different developmental stages. All sections were probed with the Flk-1 antisense probe. (A) Sagittal section of the telencephalon of a day 11.5 mouse embryo. A single blood vessel expressing Flk-1, which sprouts from the 10 meninges into the neuroectoderm, is indicated by an arrow. (B) Sagittal sections of the brain of embryo day 14.5 and (C) of postnatal day 4. Shown are regions of the mesencephalon. Branching capillaries and blood vessels expressing Flk-1 are indicated by an arrow. 15 (D) Sagittal section of an adult brain; a region of the mesencephalon is shown. Cells expressing Flk-1 are indicated by an arrow. Abbreviations: M, meninges; V, ventricle;
- FIG. 6. Expression of Flk-1 in the choroid plexus

 20 of adult brain. (A) Darkfield illumination of the
 choroid plexus of an adult mouse brain hybridized with
 Flk-1 antisense probe. (B) Choroid plexus shown at a
 higher magnification. Arrows indicate single cells,
 which show strong expression of Flk-1. Abbreviations:

 25 CP, choroid plexus; E, ependyme; Ep, epithelial cells; V,
 ventricle.
 - FIG. 7. Flk-1 is expressed in the glomeruli of the kidney. (A) Parasagittal section of a 4-day postnatal kidney, hybridized with the Flk-1 antisense probe.
- 30 Hybridization signal accumulates in the glomeruli, as indicated by arrowheads. (B) Control hybridization of an adjacent section with the sense probe. (C) Sagittal section of an adult kidney probed with Flk-1. Arrowheads indicate glomeruli. (D) Glomerulus of an adult kidney at
- 35 a higher magnification. The arrows in (A) and (D)

indicate cells aligned in strands in the juxtaglomerular region expressing Flk-1.

- FIG. 8. In situ hybridization analysis of Flk-1 expression in early embryos and extraembryonic tissues.
- 5 (A) Sagittal section of a day 8.5 mouse embryo in the maternal deciduum probed with Flk-1. (B) Higher magnification of the deciduum. Arrowheads indicate the endothelium of maternal blood vessels strongly expressing Flk-1 RNA. (C) High magnification of the yolk sac and
- 10 the trophectoderm of a day 9.5 mouse embryo. (D) High magnification of a blood island. Abbreviations:
 A, allantois; Bi, blood island; Bv, maternal blood vessel; D, deciduum; En, endodermal layer of yolk sac; M, mesenchyme; Ms, mesodermal layer of yolk sac; NF,
- 15 neural fold; T, trophoblast; Y, yolk sac.
 - FIG. 9. Flk-1 is a receptor for VEGF. (A) Cross linking of ¹²⁵I-VEGF to COS cells transiently expressing the Flk-1 receptor and control cells were incubated with ¹²⁵I-VEGF at 4°C overnight, then washed twice with
- phosphate buffered saline (PBS) and exposed to 0.5 mM of the cross linking agent DSS in PBS for 1 hour at 4°C. The cells were lysed, Flk-1 receptor immunoprecipitated, and analyzed by polyacrylamide gel electrophoresis followed by autoradiography. Molecular size markers are
- indicated in kilodaltons. (B) Specific binding of 1251-VEGF to COS cells expressing Flk-1. COS cells transiently expressing Flk-1 were removed from the plate and resuspended in binding medium (DMEM, 25 mM Hepes, 0.15% gelatin). Binding was performed at 15°C for 90
- minutes in a total volume of 0.5 ml containing 2x10⁵ cells, 15,000 cpm ¹²⁵I-VEGF, and the indicated concentrations of unlabeled ligand. The cells were washed twice with PBS / 0.1% BSA and counted in a gamma counter.

FIG. 10. VEGF-induced autophosphorylation of Flk-1. COS cells transiently expressing Flk-1 receptor and control cells were starved for 24 hours in DMEM containing 0.5% fetal calf serum and then stimulated with 5 VEGF for 10 minutes as indicated. The cells were solubilized, Flk-1 receptor immunoprecipitated with a polyclonal antibody against its C-terminus, separated by polyacrylamide gel electrophoresis, and transferred to nitrocellulose. The blot was probed with 10 antiphosphotyrosine antibodies (5B2). The protein bands

were visualized by using a horseradish-peroxidase coupled secondary antibody and BCL^{m} (Amersham) detection assay.

FIG. 11. Nucleotide Sequence of Murine Flk-1.

FIG. 12. Plasmid Maps of retroviral vector 15 constructs. pLXSN Flk-1 TM Cl.1 and pLXSN Flk-1 TM cl.3 contain Flk-1 amino acids 1 through 806. pNTK-cfms-TM contains the 541 N-terminal amino acids of c-fms.

FIG. 13. Inhibition of C6 glioblastoma tumor growth by transdominant-negative inhibition of Flk-1. C6 cells 20 were implanted either alone or coimplanted with virusproducing cells. Cell numbers are as indicated in each panel. Two different virus-producing cells lines were used: one expressing the Flk-1 TM (transdominantnegative) mutant and one expressing a transdominant-25 negative c-fms mutant (c-fms TM) as a control. Beginning at the time when the first tumors appeared, tumor volumes were measured every 2 to 3 days to obtain a growth curve. Each group is represented by four mice.

FIG. 14. Inhibition of C6 glioblastoma tumor growth 30 by transdominant-negative inhibition of Flk-1. C6 cells were implanted either alone or coimplanted with virusproducing cells. Cell numbers are as indicated in each panel. Two different virus-producing cell lines were used: one expressing the Flk-1 TM (transdominant-35 negative) mutant and one expressing a transdominantnegative c-fms mutant (cfms TM) as a control. Beginning at the time when the first tumor appeared, tumor volumes were measured every 2 to 3 days to obtain growth curve. Each group is represented by four mice.

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5. DETAILED DESCRIPTION OF THE INVENTION

The present invention relates to the use of ligands for the FLK-1 receptor to modulate angiogenesis and/or vasculogenesis. The invention also involves the

10 expression of Flk-1 to evaluate and screen for drugs and analogs of VEGF that may be involved in receptor activation, regulation and uncoupling. Such regulators of Flk-1 may be used therapeutically. For example, agonists of VEGF may be used in processes such as wound healing; in contrast, antagonists of VEGF may be used in the treatment of tumors that rely on vascularization for growth.

The invention, is based, in part, on results from in situ-hybridization and Northern blot analyses indicating
that Flk-1 is an endothelial cell specific RTK. In addition, cross-linking experiments have shown Flk-1 to be a high affinity receptor for vascular endothelial growth factor (VEGF), indicating that Flk-1 plays a crucial role in the development and differentiation of hemangioblast and in subsequent endothelial cell growth during vasculogenesis and angiogenesis.

The invention is based, also, on the discovery that expression of a transdominant-negative mutant form of the Flk-1 molecule can inhibit the biological activity of the endogenous wild type Flk-1. Experiments are descirbed herein, in which tumor cells and cells producing a recombinant retrovirus encoding a truncated Flk-1 receptor were mixed and injected into mice. Inhibition of vasculogenesis and growth of the injected tumor cells was observed in mice expressing the trucated form of the

Flk-1 receptor. Expression of transdominant negative forms of the Flk-1 molecule may be useful for treatment of diseases resulting from abnormal proliferation of blood vessels, such as rheumatoid arthritis,

5 retinopathies and growth of solid tumors.

As explained in the working examples, <u>infra</u>, the polymerase chain reaction (PCR) method was used to isolate new receptor tyrosine kinases specifically expressed in post-implantation embryos and endothelial cells. One such clone was found to encode a RTK that had almost identical sequence homology with the previously identified cDNA clone isolated from populations of cells enriched for hematopoietic cells and designated fetal liver kinase-1 (Flk-1) (Matthews et al., 1991, Proc. Natl. Acad Sci. U.S.A. 88:9026-9030) (FIG. 11).

For clarity of discussion, the invention is described in the subsections below by way of example for the murine Flk-1. However, the principles may be analogously applied to clone and express the Flk-1 of other species including humans.

5.1. THE Flk-1 CODING SEQUENCE

The nucleotide coding sequence and deduced amino acid sequence of the murine Flk-1 gene is depicted in

25 Figure 11 (SEQ. ID NO. 1) and has recently been described in Matthews et al., 1991, Proc. Natl. Acad. Sci. U.S.A., 88:9026-9030. In accordance with the invention, the nucleotide sequence of the Flk-1 protein or its functional equivalent in mammals, including humans, can be used to generate recombinant molecules which direct the expression of Flk-1; hereinafter, this receptor will be referred to as "Flk-1", regardless of the species from which it is derived.

In a specific embodiment described herein, the 35 murine Flk-1 gene was isolated by performing a polymerase

chain reaction (PCR) using two degenerate oligonucleotide primer pools that were designed on the basis of highly conserved sequences within the kinase domain of receptor tyrosine kinases (Hanks et al., 1988,) As a template,

5 DNA from a \(\lambda \)gt10 cDNA library prepared from day 8.5 mouse embryos, was used. In a parallel approach, similar primers were used to amplify RTK cDNA sequences from capillary endothelial cells that had been isolated from the brains of post-natal day 4-8 mice. This is a time

10 when brain endothelial cell proliferation is maximal. Both approaches yielded cDNA sequences encoding the recently described fetal liver RTK, Flk-1 (Matthews et al., 1991). Based on amino acid homology, this receptor is a member of the type III subclass of RTKs (Ullrich and Schlessinger) which contain immunoglobulin-like repeats

The invention also relates to Flk-1 genes isolated from other species, including humans, in which Flk-1 activity exists. Members of the Flk-1 family are defined 20 herein as those receptors that bind VEGF or fragments of the peptide. Such receptors may demonstrate about 80% homology at the amino acid level in substantial stretches of DNA sequence. A bacteriophage cDNA library may be screened, under conditions of reduced stringency, using a 25 radioactively labeled fragment of the mouse Flk-1 clone. Alternatively the mouse Flk-1 sequence can be used to design degenerate or fully degenerate oligonucleotide probes which can be used as PCR probes or to screen bacteriophage cDNA libraries. A polymerase chain 30 reaction (PCR) based strategy may be used to clone human Flk-1. Two pools of degenerate oligonucleotides, corresponding to a conserved motifs between the mouse Flk-1 and receptor tyrosine kinases, may be designed to serve as primers in a PCR reaction. The template for the

35 reaction is cDNA obtained by reverse transcription of

in their extracellular domains (FIG. 1).

mRNA prepared from cell lines or tissue known to express human Flk-1. The PCR product may be subcloned and sequenced to insure that the amplified sequences represent the Flk-1 sequences. The PCR fragment may be used to isolate a full length Flk-1 cDNA clone by radioactively labeling the amplified fragment and screening a bacteriophage cDNA library. Alternatively, the labeled fragment may be used to screen a genomic library. For a review of cloning strategies which may be used, see e.g., Maniatis, 1989, Molecular Cloning, A Laboratory Manual, Cold Springs Harbor Press, N.Y.; and Ausubel et al., 1989, Current Protocols in Molecular Biology, (Green Publishing Associates and Wiley Interscience, N.Y.)

Isolation of a human Flk-1 cDNA may also be achieved by construction of a cDNA library in a mammalian expression vector such as pcDNA1, that contains SV40 origin of replication sequences which permit high copy number expression of plasmids when transferred into COS cells. The expression of Flk-1 on the surface of transfected COS cells may be detected in a number of ways, including the use of a labeled ligand such as VEGF or a VEGF agonist labeled with a radiolabel, fluorescent label or an enzyme. Cells expressing the human Flk-1 may be enriched by subjecting transfected cells to a FACS (fluorescent activated cell sorter) sort.

In accordance with the invention, Flk-1 nucleotide sequences which encode Flk-1, peptide fragments of Flk-1, Flk-1 fusion proteins or functional equivalents thereof

may be used to generate recombinant DNA molecules that direct the expression of Flk-1 protein or a functionally equivalent thereof, in appropriate host cells.

Alternatively, nucleotide sequences which hybridize to portions of the Flk-1 sequence may also be used in

nucleic acid hybridization assays, Southern and Northern blot analyses, etc.

Due to the inherent degeneracy of the genetic code, other DNA sequences which encode substantially the same 5 or a functionally equivalent amino acid sequence, may be used in the practice of the invention for the cloning and expression of the Flk-1 protein. Such DNA sequences include those which are capable of hybridizing to the murine Flk-1 sequence under stringent conditions.

Altered DNA sequences which may be used in accordance with the invention include deletions, additions or substitutions of different nucleotide residues resulting in a sequence that encodes the same or a functionally equivalent gene product. The gene product itself may 15 contain deletions, additions or substitutions of amino acid residues within the Flk-1 sequence, which result in a silent change thus producing a functionally equivalent Flk-1. Such amino acid substitutions may be made on the basis of similarity in polarity, charge, solubility, 20 hydrophobicity, hydrophilicity, and/or the amphipatic

nature of the residues involved. For example, negatively charged amino acids include aspartic acid and glutamic acid; positively charged amino acids include lysine and arginine; amino acids with uncharged polar head groups 25 having similar hydrophilicity values include the

following: leucine, isoleucine, valine; glycine, analine; asparagine, glutamine; serine, threonine; phenylalanine, tyrosine. As used herein, a functionally equivalent Flk-1 refers to a receptor which binds to VEGF or fragments,

30 but not necessarily with the same binding affinity of its counterpart native Flk-1.

The DNA sequences of the invention may be engineered in order to alter the Flk-1 coding sequence for a variety of ends including but not limited to alterations which 35 modify processing and expression of the gene product.

For example, mutations may be introduced using techniques which are well known in the art, e.g. site-directed mutagenesis, to insert new restriction sites, to alter glycosylation patterns, phosphorylation, etc. For example, in certain expression systems such as yeast, host cells may over glycosylate the gene product. When using such expression systems it may be preferable to alter the Flk-1 coding sequence to eliminate any N-linked glycosylation site.

In another embodiment of the invention, the Flk-1 or a modified Flk-1 sequence may be ligated to a heterologous sequence to encode a fusion protein. For example, for screening of peptide libraries it may be useful to encode a chimeric Flk-1 protein expressing a heterologous epitope that is recognized by a commercially available antibody. A fusion protein may also be engineered to contain a cleavage site located between the Flk-1 sequence and the heterologous protein sequence, so that the Flk-1 can be cleaved away from the heterologous moiety.

In an alternate embodiment of the invention, the coding sequence of Flk-1 could be synthesized in whole or in part, using chemical methods well known in the art. See, for example, Caruthers, et al., 1980, Nuc. Acids Res. Symp. Ser. 7:215-233; Crea and Horn, 180, Nuc. Acids Res. 9(10):2331; Matteucci and Caruthers, 1980, Tetrahedron Letters 21:719; and Chow and Kempe, 1981, Nuc. Acids Res. 9(12):2807-2817. Alternatively, the protein itself could be produced using chemical methods to synthesize the Flk-1 amino acid sequence in whole or in part. For example, peptides can be synthesized by solid phase techniques, cleaved from the resin, and purified by preparative high performance liquid chromatography. (E.g., see Creighton, 1983, Proteins Structures And Molecular Principles, W.H. Freeman and

Co., N.Y. pp. 50-60). The composition of the synthetic peptides may be confirmed by amino acid analysis or sequencing (e.g., the Edman degradation procedure; see Creighton, 1983, Proteins, Structures and Molecular Principles, W.H. Freeman and Co., N.Y., pp. 34-49.

5.2. EXPRESSION OF Flk-1 RECEPTOR AND GENERATION OF CELL LINES THAT EXPRESS Flk-1

In order to express a biologically active Flk-1, the

nucleotide sequence coding for Flk-1, or a functional
equivalent as described in Section 5.1 supra, is inserted
into an appropriate expression vector, i.e., a vector
which contains the necessary elements for the
transcription and translation of the inserted coding

sequence. The Flk-1 gene products as well as host cells
or cell lines transfected or transformed with recombinant
Flk-1 expression vectors can be used for a variety of
purposes. These include but are not limited to
generating antibodies (i.e., monoclonal or polyclonal)

that bind to the receptor, including those that
competitively inhibit binding of VEGF and "neutralize"
activity of Flk-1 and the screening and selection of VEGF
analogs or drugs that act via the Flk-1 receptor; etc.

5.2.1. EXPRESSION SYSTEMS

Methods which are well known to those skilled in the art can be used to construct expression vectors containing the Flk-1 coding sequence and appropriate transcriptional/translational control signals. These methods include in vitro recombinant DNA techniques, synthetic techniques and in vivo recombination/genetic recombination. See, for example, the techniques described in Maniatis et al., 1989, Molecular Cloning A Laboratory Manual, Cold Spring Harbor Laboratory, N.Y. and Ausubel et al., 1989, Current Protocols in Molecular

Biology, Greene Publishing Associates and Wiley Interscience, N.Y.

A variety of host-expression vector systems may be utilized to express the Flk-1 coding sequence. These 5 include but are not limited to microorganisms such as bacteria transformed with recombinant bacteriophage DNA, plasmid DNA or cosmid DNA expression vectors containing the Flk-1 coding sequence; yeast transformed with recombinant yeast expression vectors containing the Flk-1 10 coding sequence; insect cell systems infected with recombinant virus expression vectors (e.g., baculovirus) containing the Flk-1 coding sequence; plant cell systems infected with recombinant virus expression vectors (e.g., cauliflower mosaic virus, CaMV; tobacco mosaic virus, 15 TMV) or transformed with recombinant plasmid expression vectors (e.g., Ti plasmid) containing the Flk-1 coding sequence; or animal cell systems infacted with recombinant virus expression vectors (e.g., adenovirus, vaccinia virus) including cell lines engineered to 20 contain multiple copies of the Flk-1 DNA either stably amplified (CHO/dhfr) or unstably amplified in double-

minute chromosomes (e.g., murine cell lines).
The expression elements of these systems vary in their strength and specificities. Depending on the host/vector system utilized, any of a number of suitable transcription and translation elements, including constitutive and inducible promoters, may be used in the expression vector. For example, when cloning in bacterial systems, inducible promoters such as pL of bacteriophage λ, plac, ptrp, ptac (ptrp-lac hybrid promoter) and the like may be used; when cloning in insect cell systems, promoters such as the baculovirus polyhedrin promoter may be used; when cloning in plant

cell systems, promoters derived from the genome of plant cells (e.g., heat shock promoters; the promoter for the

small subunit of RUBISCO; the promoter for the chlorophyll a/b binding protein) or from plant viruses (e.g., the 35S RNA promoter of CaMV; the coat protein promoter of TMV) may be used; when cloning in mammalian cell systems, promoters derived from the genome of mammalian cells (e.g., metallothionein promoter) or from mammalian viruses (e.g., the adenovirus late promoter; the vaccinia virus 7.5K promoter) may be used; when generating cell lines that contain multiple copies of the Flk-1 DNA SV40-, BPV- and EBV-based vectors may be used with an appropriate selectable marker.

In bacterial systems a number of expression vectors may be advantageously selected depending upon the use intended for the Flk-1 expressed. For example, when 15 large quantities of Flk-1 are to be produced for the generation of antibodies or to screen peptide libraries, vectors which direct the expression of high levels of fusion protein products that are readily purified may be desirable. Such vectors include but are not limited to 20 the E. coli expression vector pUR278 (Ruther et al., 1983, EMBO J. 2:1791), in which the Flk-1 coding sequence may be ligated into the vector in frame with the lac Z coding region so that a hybrid AS-lac Z protein is produced; pIN vectors (Inouye & Inouye, 1985, Nucleic 25 acids Res. 13:3101-3109; Van Heeke & Schuster, 1989, J. Biol. Chem. 264:5503-5509); and the like. pGEX vectors may also be used to express foreign polypeptides as fusion proteins with glutathione S-transferase (GST). In general, such fusion proteins are soluble and can easily 30 be purified from lysed cells by adscrption to glutathione-agarose beads followed by elution in the presence of free glutathione. The pGEX vectors are designed to include thrombin or factor Xa protease cleavage sites so that the cloned polypeptide of interest 35 can be released from the GST moiety.

In yeast, a number of vectors containing constitutive or inducible promoters may be used. For a review see, Current Protocols in Molecular Biology, Vol. 2, 1988, Ed. Ausubel et al., Greene Publish. Assoc. & Wiley Interscience, Ch. 13; Grant et al., 1987, Expression and Secretion Vectors for Yeast, in Methods in Enzymology, Eds. Wu & Grossman, 1987, Acad. Press, N.Y., Vol. 153, pp. 516-544; Glover, 1986, DNA Cloning, Vol. II, IRL Press, Wash., D.C., Ch. 3; and Bitter, 1987, Heterologous Gene Expression in Yeast, Methods in Enzymology, Eds. Berger & Kimmel, Acad. Press, N.Y., Vol. 152, pp. 673-684; and The Molecular Biology of the Yeast Saccharomyces, 1982, Eds. Strathern et al., Cold Spring Harbor Press, Vols. I and II.

In cases where plant expression vectors are used, 15 the expression of the Flk-1 coding sequence may be driven by any of a number of promoters. For example, viral promoters such as the 35S RNA and 19S RNA promoters of CaMV (Brisson et al., 1984, Nature 310:511-514), or the 20 coat protein promoter of TMV (Takamatsu et al., 1987, EMBO J. 6:307-311) may be used; alternatively, plant promoters such as the small subunit of RUBISCO (Coruzzi et al., 1984, EMBO J. 3:1671-1680; Broglie et al., 1984, Science 224:838-843); or heat shock promoters, e.g., 25 soybean hsp17.5-E or hsp17.3-B (Gurley et al., 1986, Mol. Cell. Biol. 6:559-565) may be used. These constructs can be introduced into plant cells using Ti plasmids, Ri plasmids, plant virus vectors, direct DNA transformation, microinjection, electroporation, etc. For reviews of 30 such techniques see, for example, Weissbach & Weissbach, 1988, Methods for Plant Molecular Biology, Academic Press, NY, Section VIII, pp. 421-463; and Grierson & Corey, 1988, Plant Molecular Biology, 2d Ed., Blackie, London, Ch. 7-9.

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An alternative expression system which could be used to express Flk-1 is an insect system. In one such system, Autographa californica nuclear polyhidrosis virus (AcNPV) is used as a vector to express foreign genes.

- 5 The virus grows in Spodoptera frugiperda cells. The Flk-1 coding sequence may be cloned into non-essential regions (for example the polyhedrin gene) of the virus and placed under control of an AcNPV promoter (for example the polyhedrin promoter). Successful insertion
- 10 of the Flk-1 coding sequence will result in inactivation of the polyhedrin gene and production of non-occluded recombinant virus (i.e., virus lacking the proteinaceous coat coded for by the polyhedrin gene). These recombinant viruses are then used to infect Spodoptera
- 15 frugiperda cells in which the inserted gene is expressed. (E.g., see Smith et al., 1983, J. Viol. 46:584; Smith, U.S. Patent No. 4,215,051).

In mammalian host cells, a number of viral based expression systems may be utilized. In cases where an 20 adenovirus is used as an expression vector, the Flk-1 coding sequence may be ligated to an adenovirus transcription/translation control complex, e.g., the late promoter and tripartite leader sequence. This chimeric gene may then be inserted in the adenovirus genome by in 25 vitro or in vivo recombination. Insertion in a nonessential region of the viral genome (e.g., region El or E3) will result in a recombinant virus that is viable and

capable of expressing Flk-1 in infected hosts. See Logan & Shenk, 1984, Proc. Natl. Acad. Sci. (USA) 30 81:3655-3659). Alternatively, the vaccinia 7.5K promoter may be used. (See, e.g., Mackett et al., 1982, Proc. Natl. Acad. Sci. (USA) 79:7415-7419; Mackett et al., 1984, J. Virol. 49:857-864; Panicali et al., 1982, Proc.

(E.q.,

Natl. Acad. Sci. 79:4927-4931).

Specific initiation signals may also be required for efficient translation of inserted Flk-1 coding sequences. These signals include the ATG initiation codon and adjacent sequences. In cases where the entire Flk-1 gene, including its own initiation codon and adjacent sequences, is inserted into the appropriate expression vector, no additional translational control signals may be needed. However, in cases where only a portion of the Flk-1 coding sequence is inserted, exogenous

translational control signals, including the ATG initiation codon, must be provided. Furthermore, the initiation codon must be in phase with the reading frame of the Flk-1 coding sequence to ensure translation of the entire insert. These exogenous translational control

signals and initiation codons can be of a variety of origins, both natural and synthetic. The efficiency of expression may be enhanced by the inclusion of appropriate transcription enhancer elements, transcription terminators, etc. (see Bittner et al., 1987, Methods in Enzymol. 153:516-544).

In addition, a host cell strain may be chosen which modulates the expression of the inserted sequences, or modifies and processes the gene product in the specific fashion desired. Such modifications (e.g.,

glycosylation) and processing (e.g., cleavage) of protein products may be important for the function of the protein. Different host cells have characteristic and specific mechanisms for the post-translational processing and modification of proteins. Appropriate cells lines or

nodification and processing of the foreign protein expressed. To this end, eukaryotic host cells which possess the cellular machinery for proper processing of the primary transcript, glycosylation, and

35 phosphorylation of the gene product may be used. Such

mammalian host cells include but are not limited to CHO, VERO, BHK, HeLa, COS, MDCK, 293, WI38, etc.

For long-term, high-yield production of recombinant proteins, stable expression is preferred. For example, 5 cell lines which stably express the Flk-1 may be engineered. Rather than using expression vectors which contain viral origins of replication, host cells can be transformed with the Flk-1 DNA controlled by appropriate expression control elements (e.g., promoter, enhancer, 10 sequences, transcription terminators, polyadenylation sites, etc.), and a selectable marker. Following the introduction of foreign DNA, engineered cells may be allowed to grow for 1-2 days in an enriched media, and then are switched to a selective media. The selectable 15 marker in the recombinant plasmid confers resistance to the selection and allows cells to stably integrate the plasmid into their chromosomes and grow to form foci which in turn can be cloned and expanded into cell lines. This method may advantageously be used to engineer cell 20 lines which express the Flk-1 on the cell surface, and which respond to VEGF mediated signal transduction. Such engineered cell lines are particularly useful in screening VEGF analogs.

A number of selection systems may be used, including

but not limited to the herpes simplex virus thymidine

kinase (Wigler, et al., 1977, Cell 11:223), hypoxanthine
guanine phosphoribosyltransferase (Szybalska & Szybalski,

1962, Proc. Natl. Acad. Sci. USA 48:2026), and adenine

phosphoribosyltransferase (Lowy, et al., 1980, Cell

22:817) genes can be employed in tk, hgprt or aprt

cells, respectively. Also, antimetabolite resistance can

be used as the basis of selection for dhfr, which confers

resistance to methotrexate (Wigler, et al., 1980, Natl.

Acad. Sci. USA 77:3567; O'Hare, et al., 1981, Proc. Natl.

35 Acad. Sci. USA 78:1527); gpt, which confers resistance to

mycophenolic acid (Mulligan & Berg, 1981), Proc. Natl.
Acad. Sci. USA 78:2072); neo, which confers resistance to
the aminoglycoside G-418 (Colberre-Garapin, et al., 1981,
J. Mol. Biol. 150:1); and hygro, which confers resistance
to hygromycin (Santerre, et al., 1984, Gene 30:147)
genes. Recently, additional selectable genes have been
described, namely trpB, which allows cells to utilize
indole in place of tryptophan; hisD, which allows cells
to utilize histinol in place of histidine (Hartman &
Mulligan, 1988, Proc. Natl. Acad. Sci. USA 85:8047); and
ODC (ornithine decarboxylase) which confers resistance to
the ornithine decarboxylase inhibitor, 2(difluoromethyl)-DL-ornithine, DFMO (McConloque L., 1987,
In: Current Communications in Molecular Biology, Cold
Spring Harbor Laboratory ed.).

5.2.2. IDENTIFICATION OF TRANSFECTANTS OR TRANSFORMANTS THAT EXPRESS THE F1k-1

The host cells which contain the coding sequence and which express the biologically active gene product may be identified by at least four general approaches; (a) DNA-DNA or DNA-RNA hybridization; (b) the presence or absence of "marker" gene functions; (c) assessing the level of transcription as measured by the expression of Flk-1 mRNA transcripts in the host cell; and (d) detection of the gene product as measured by immunoassay or by its biological activity.

In the first approach, the presence of the Flk-1 coding sequence inserted in the expression vector can be detected by DNA-DNA or DNA-RNA hybridization using probes comprising nucleotide sequences that are homologous to the Flk-1 coding sequence, respectively, or portions or derivatives thereof.

In the second approach, the recombinant expression 35 vector/host system can be identified and selected based upon the presence or absence of certain "marker" gene functions (e.g., thymidine kinase activity, resistance to antibiotics, resistance to methotrexate, transformation phenotype, occlusion body formation in baculovirus,

5 etc.). For example, if the Flk-1 coding sequence is inserted within a marker gene sequence of the vector, recombinants containing the Flk-1 coding sequence can be identified by the absence of the marker gene function.

Alternatively, a marker gene can be placed in tandem with the Flk-1 sequence under the control of the same or different promoter used to control the expression of the Flk-1 coding sequence. Expression of the marker in response to induction or selection indicates expression

of the Flk-1 coding sequence.

In the third approach, transcriptional activity for the Flk-1 coding region can be assessed by hybridization assays. For example, RNA can be isolated and analyzed by Northern blot using a probe homologous to the Flk-1 coding sequence or particular portions thereof.

20 Alternatively, total nucleic acids of the host cell may be extracted and assayed for hybridization to such probes.

In the fourth approach, the expression of the Flk-1 protein product can be assessed immunologically, for example by Western blots, immunoassays such as radioimmuno-precipitation, enzyme-linked immunoassays and the like. The ultimate test of the success of the expression system, however, involves the detection of the biologically active Flk-1 gene product. A number of assays can be used to detect receptor activity including but not limited to VEGF binding assays; and VEGF biological assays using engineered cell lines as the test substrate.

5.3. USES OF THE F1k-1 RECEPTOR AND ENGINEERED CELL LINES

Angiogenesis, the growth of new blood capillary
vessels, is required for a number of physiological

5 processes ranging from wound healing, tissue and organ
regeneration, placental formation after pregnancy and
embryonic development. Abnormal proliferation of blood
vessels is an important component of a variety of
diseases such as rheumatoid arthritis, retinopathies, and
10 psoriasis. Angiogenesis is also an important factor in
the growth and metastatic activity of solid tumors that
rely on vascularization. Therefore, inhibitors of
angiogenesis may be used therapeutically for the
treatment of diseases resulting from or accompanied by
abnormal growth of blood vessels and for treatments of
malignancies involving growth and spread of solid tumors.

In an embodiment of the invention the Flk-1 receptor and/or cell lines that express the Flk-1 receptor may be used to screen for antibodies, peptides, or other ligands that act as agonists or antagonists of angiogenesis or vasculogenesis mediated by the Flk-1 receptor. For example, anti-Flk-1 antibodies capable of neutralizing the activity of VEGF, may be used to inhibit Flk-1 function. Additionally, anti-Flk-1 antibodies which mimic VEGF activity may be selected for uses in wound healing. Alternatively, screening of peptide libraries with recombinantly expressed soluble Flk-1 protein or cell lines expressing Flk-1 protein may be useful for identification of therapeutic molecules that function by inhibiting the biological activity of Flk-1.

In an embodiment of the invention, engineered cell lines which express the entire Flk-1 coding region or its ligand binding domain may be utilized to screen and identify VEGF antagonists as well as agonists. Synthetic compounds, natural products, and other sources of

potentially biologically active materials can be screened in a number of ways. The ability of a test compound to inhibit binding of VEGF to Flk-1 may be measured using standard receptor binding techniques, such as those described in Section 6.1.9. The ability of agents to prevent or mimic, the effect of VEGF binding on signal transduction responses on Flk-1 expressing cells may be measured. For example, responses such as activation of Flk-1 kinase activity, modulation of second messenger production or changes in cellular metabolism may be monitored. These assays may be performed using conventional techniques developed for these purposes.

5.3.1. SCREENING OF PEPTIDE LIBRARY WITH F1k-1 PROTEIN OR ENGINEERED CELL LINES

Random peptide libraries consisting of all possible combinations of amino acids attached to a solid phase support may be used to identify peptides that are able to bind to the ligand binding site of a given receptor or other functional domains of a receptor such as kinase domains (Lam, K.S. et al., 1991, Nature 354: 82-84). The screening of peptide libraries may have therapeutic value in the discovery of pharmaceutical agents that act to inhibit the biological activity of receptors through their interactions with the given receptor.

Identification of molecules that are able to bind to the Flk-1 may be accomplished by screening a peptide library with recombinant soluble Flk-1 protein. Methods for expression and purification of Flk-1 are described in Section 5.2.1 and may be used to express recombinant full length Flk-1 or fragments of Flk-1 depending on the functional domains of interest. For example, the kinase and extracellular ligand binding domains of Flk-1 may be separately expressed and used to screen peptide

35 libraries.

To identify and isolate the peptide/solid phase support that interacts and forms a complex with Flk-1, it is necessary to label or "tag" the Flk-1 molecule. The Flk-1 protein may be conjugated to enzymes such as 5 alkaline phosphatase or horseradish peroxidase or to other reagents such as fluorescent labels which may include fluorescein isothyiocynate (FITC), phycoerythrin (PE) or rhodamine. Conjugation of any given label, to Flk-1, may be performed using techniques that are routine 10 in the art. Alternatively, Flk-1 expression vectors may be engineered to express a chimeric Flk-1 protein containing an epitope for which a commercially available antibody exist. The epitope specific antibody may be tagged using methods well known in the art including 15 labeling with enzymes, fluorescent dyes or colored or magnetic beads.

The "tagged" Flk-1 conjugate is incubated with the random peptide library for 30 minutes to one hour at 22°C to allow complex formation between Flk-1 and peptide 20 species within the library. The library is then washed to remove any unbound Flk-1 protein. If Flk-1 has been conjugated to alkaline phosphatase or horseradish peroxidase the whole library is poured into a petri dish containing a substrates for either alkaline phosphatase 25 or peroxidase, for example, 5-bromo-4-chloro-3-indoyl phosphate (BCIP) or 3,3',4,4"-diamnobenzidine (DAB), respectively. After incubating for several minutes, the peptide/solid phase-Flk-1 complex changes color, and can be easily identified and isolated physically under a 30 dissecting microscope with a micromanipulator. If a fluorescent tagged Flk-1 molecule has been used, complexes may be isolated by fluorescent activated sorting. If a chimeric Flk-1 protein expressing a heterologous epitope has been used, detection of the 35 peptide/Flk-1 complex may be accomplished by using a

labeled epitope specific antibody. Once isolated, the identity of the peptide attached to the solid phase support may be determined by peptide sequencing.

In addition to using soluble Flk-1 molecules, in 5 another embodiment, it is possible to detect peptides that bind to cell surface receptors using intact cells. The use of intact cells is preferred for use with receptors that are multi-subunits or labile or with receptors that require the lipid domain of the cell 10 membrane to be functional. Methods for generating cell lines expressing Flk-1 are described in Sections 5.2.1. and 5.2.2. The cells used in this technique may be either live or fixed cells. The cells will be incubated with the random peptide library and will bind to certain 15 peptides in the library to form a "rosette" between the target cells and the relevant solid phase support/peptide. The rosette can thereafter be isolated by differential centrifugation or removed physically under a dissecting microscope.

As an alternative to whole cell assays for membrane bound receptors or receptors that require the lipid domain of the cell membrane to be functional, the receptor molecules can be reconstituted into liposomes where label or "tag" can be attached.

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5.3.2. ANTIBODY PRODUCTION AND SCREENING

Various procedures known in the art may be used for the production of antibodies to epitopes of the recombinantly produced Flk-1 receptor. Such antibodies include but are not limited to polyclonal, monoclonal, chimeric, single chain, Fab fragments and fragments produced by an Fab expression library. Neutralizing antibodies <u>i.e.</u>, those which compete for the VEGF binding

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site of the receptor are especially preferred for diagnostics and therapeutics.

Monoclonal antibodies that bind Flk-1 may be radioactively labeled allowing one to follow their location and distribution in the body after injection. Radioactivity tagged antibodies may be used as a non-invasive diagnostic tool for imaging de novo vascularization associated with a number of diseases including rheumatoid arthritis, macular degeneration, and formation of tumors and metastases.

Immunotoxins may also be designed which target cytotoxic agents to specific sites in the body. For example, high affinity Flk-1 specific monoclonal antibodies may be covalently complexed to bacterial or plant toxins, such as diptheria toxin, abrin or ricin. A general method of preparation of antibody/hybrid molecules may involve use of thiol-crosslinking reagents such as SPDP, which attack the primary amino groups on the antibody and by disulfide exchange, attach the toxin to the antibody. The hybrid antibodies may be used to specifically eliminate Flk-1 expressing endothelial cells.

For the production of antibodies, various host animals may be immunized by injection with the Flk-1

25 protein including but not limited to rabbits, mice, rats, etc. Various adjuvants may be used to increase the immunological response, depending on the host species, including but not limited to Freund's (complete and incomplete), mineral gels such as aluminum hydroxide,

30 surface active substances such as lysolecithin, pluronic polyols, polyanions, peptides, oil emulsions, keyhole limpet hemocyanin, dinitrophenol, and potentially useful human adjuvants such as BCG (bacille Calmette-Guerin) and Corvnebacterium parvum.

Monoclonal antibodies to Flk-1 may be prepared by using any technique which provides for the production of antibody molecules by continuous cell lines in culture. These include but are not limited to the hybridoma

- 5 technique originally described by Kohler and Milstein, (Nature, 1975, 256:495-497), the human B-cell hybridoma technique (Kosbor et al., 1983, Immunology Today, 4:72; Cote et al., 1983, Proc. Natl. Acad. Sci., 80:2026-2030) and the EBV-hybridoma technique (Cole et al., 1985,
- Monoclonal Antibodies and Cancer Therapy, Alan R. Liss, Inc., pp. 77-96). In addition, techniques developed for the production of "chimeric antibodies" (Morrison et al., 1984, Proc. Natl. Acad. Sci., 81:6851-6855; Neuberger et al., 1984, Nature, 312:604-608; Takeda et al., 1985,
- Nature, 314:452-454) by splicing the genes from a mouse antibody molecule of appropriate antigen specificity together with genes from a human antibody molecule of appropriate biological activity can be used.

 Alternatively, techniques described for the production of

20 single chain antibodies (U.S. Patent 4,946,778) can be adapted to produce Flk-1-specific single chain antibodies.

Antibody fragments which contain specific binding sites of Flk-1 may be generated by known techniques. For example, such fragments include but are not limited to: the F(ab')₂ fragments which can be produced by pepsin digestion of the antibody molecule and the Fab fragments which can be generated by reducing the disulfide bridges of the F(ab')₂ fragments. Alternatively, Fab expression libraries may be constructed (Huse et al., 1989, Science, 246:1275-1281) to allow rapid and easy identification of monoclonal Fab fragments with the desired specificity to Flk-1.

5.4. USES OF F1k-1 CODING SEQUENCE

The Flk-1 coding sequence may be used for diagnostic purposes for detection of Flk-1 expression. Included in the scope of the invention are oligoribonucleotide sequences, that include antisense RNA and DNA molecules 5 and ribozymes that function to inhibit translation of Flk-1. In addition, mutated forms of Flk-1, having a dominant negative effect, may be expressed in targeted cell populations to inhibit the activity of endogenously expressed wild-type Flk-1.

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5.4.1. USE OF Flk-1 CODING SEQUENCE IN DIAGNOSTICS AND THERAPEUTICS

The F1k-1 DNA may have a number of uses for the diagnosis of diseases resulting from aberrant expression 15 of F1k-1. For example, the F1k-1 DNA sequence may be used in hybridization assays of biopsies or autopsies to diagnose abnormalities of Flk-1 expression; e.g., Southern or Northern analysis, including in situ hybridization assays.

The Flk-1 cDNA may be used as a probe to detect the expression of the Flk-1 mRNA. In a specific example described herein, the expression of Flk-1 mRNA in mouse embryos of different developmental stages was analyzed. Northern blot analysis indicated abundant expression of a 25 major 5.5 kb mRNA between day 9.5 and day 18.5, with apparent decline towards the end of gestation (FIG. 2A). In post-natal day 4-8 brain capillaries Flk-1 mRNA was found to be highly enriched compared to total brain RNA (FIG.2B), suggesting a role for Flk-1 in endothelial cell 30 proliferation.

To obtain more detailed information about the expression of Flk-1 during embryonic development and during the early stages of vascular development in situ hybridization experiments were performed as described in 35 Section 6.1.4. In situ hybridizations demonstrated that

Flk-1 expression in vivo during embryonic mouse development is largely restricted to endothelial cells and their precursors (FIG. 3 and FIG. 4). Flk-1 is expressed in endothelial cells during physiological 5 processes that are characterized by endothelial cell proliferation and the temporal and spatial expression pattern found in the embryonic brain correlate precisely with the development of the neural vascular system as described by Bar (1980). Vascular sprouts originating in 10 the perineural plexus grow radially into the neuroectoderm and branch there and these sprouts were found to express high amounts of Flk-1 mRNA (FIG. 5). In the early postnatal stages endothelial cell proliferation is still evident and Flk-1 is expressed, whereas in the 15 adult organism, after completion of the vascularization process, the decline in endothelial cell proliferation parallels a decrease in Flk-1 expression.

Also within the scope of the invention are oligoribonucleotide sequences, that include anti-sense RNA and
DNA molecules and ribozymes that function to inhibit the
translation of Flk-1 mRNA. Anti-sense RNA and DNA
molecules act to directly block the translation of mRNA
by binding to targeted mRNA and preventing protein
translation. In regard to antisense DNA,

oligodeoxyribonucleotides derived from the translation initiation site, <u>e.g.</u>, between -10 and +10 regions of the Flk-1 nucleotide sequence, are preferred.

Ribozymes are enzymatic RNA molecules capable of catalyzing the specific cleavage of RNA. The mechanism of ribozyme action involves sequence specific hybridization of the ribozyme molecule to complementary target RNA, followed by a endonucleolytic cleavage. Within the scope of the invention are engineered hammerhead motif ribozyme molecules that specifically and

efficiently catalyze endonucleolytic cleavage of Flk-1 RNA sequences.

Specific ribozyme cleavage sites within any potential RNA target are initially identified by scanning the target molecule for ribozyme cleavage sites which include the following sequences, GUA, GUU and GUC. Once identified, short RNA sequences of between 15 and 20 ribonucleotides corresponding to the region of the target gene containing the cleavage site may be evaluated for predicted structural features such as secondary structure that may render the oligonucleotide sequence unsuitable. The suitability of candidate targets may also be evaluated by testing their accessibility to hybridization with complementary oligonucleotides, using ribonuclease protection assays.

Both anti-sense RNA and DNA molecules and ribozymes of the invention may be prepared by any method known in the art for the synthesis of RNA molecules. These include techniques for chemically synthesizing oligodeoxyribonucleotides well known in the art such as for example solid phase phosphoramidite chemical synthesis. Alternatively, RNA molecules may be generated by in vitro and in vivo transcription of DNA sequences encoding the antisense RNA molecule. Such DNA sequences may be incorporated into a wide variety of vectors which incorporate suitable RNA polymerase promoters such as the T7 or SP6 polymerase promoters. Alternatively, antisense cDNA constructs that synthesize antisense RNA constitutively or inducibly, depending on the promoter used, can be introduced stably into cell lines.

Various modifications to the DNA molecules may be introduced as a means of increasing intracellular stability and half-life. Possible modifications include but are not limited to the addition of flanking sequences of ribo- or deoxy- nucleotides to the 5' and/or 3' ends

of the molecule or the use of phosphorothicate or 2' O-methyl rather than phosphodiesterase linkages within the oligodeoxyribonucleotide backbone.

5.4.2. USE OF DOMINANT NEGATIVE Flk-1 MUTANTS IN GENE THERAPY

Receptor dimerization induced by ligands, is thought to provide an allosteric regulatory signal that functions to couple ligand binding to stimulation of kinase activity. Defective receptors can function as dominant negative mutations by suppressing the activation and response of normal receptors by formation of unproductive heterodimers. Therefore, defective receptors can be engineered into recombinant viral vectors and used in gene therapy in individuals that inappropriately express Flk-1.

In an embodiment of the invention, mutant forms of the Flk-1 molecule having a dominant negative effect may be identified by expression in selected cells. Deletion or missense mutants of Flk-1 that retain the ability to form dimers with wild type Flk-1 protein but cannot function in signal transduction may be used to inhibit the biological activity of the endogenous wild type Flk-1. For example, the cytoplasmic kinase domain of Flk-1 may be deleted resulting in a truncated Flk-1 molecule that is still able to undergo dimerization with endogenous wild type receptors but unable to transduce a signal.

Abnormal proliferation of blood vessels is an
important component of a variety of pathogenic disorders
such as rheumatoid arthritis, retinopathies and
psoriasis. Uncontrolled angiogenesis is also an
important factor in the growth and metastases of solid
tumors. Recombinant viruses may be engineered to express
dominant negative forms of Flk-1 which may be used to

inhibit the activity of the wild type endogenous Flk-1. These viruses may be used therapeutically for treatment of diseases resulting from aberrant expression or activity of Flk-1.

retroviruses, vaccinia virus, adeno-associated virus, herpes viruses, or bovine papilloma virus, may be used for delivery of recombinant Flk-1 into the targeted cell population. Methods which are well known to those skilled in the art can be used to construct recombinant viral vectors containing Flk-1 coding sequence. See, for example, the techniques described in Maniatis et al., 1989, Molecular Cloning A Laboratory Manual, Cold Spring Harbor Laboratory, N.Y. and Ausubel et al., 1989, Current Protocols in Molecular Biology, Greene Publishing Associates and Wiley Interscience, N.Y. Alternatively, recombinant Flk-1 molecules can be reconstituted into liposomes for delivery to target cells.

In a specific embodiment of the invention, a

20 deletion mutant of the Flk-1 receptor was engineered into
a recombinant retroviral vector. Two clonal isolates
designated pLXSN Flk-1 TM cl.1 and pLXSN Flk-1 TM cl.3
contain a truncated Flk-1 receptor lacking the 561 COOHterminal amino acids. To obtain virus producing cell
lines, PA37 cells were transfected with the recombinant
vectors and, subsequently, conditioned media containing
virus were used to infect GPE cells.

To test whether expression of signaling-defective mutants inhibits endogenous Flk-1 receptor activity, C6

30 rat gliobastoma cells (tumor cells) and mouse cells producing the recombinant retroviruses were mixed and injected subcutaneously into nude mice. Normally, injection of tumor cells into nude mice would result in proliferation of the tumor cells and vascularization of the resulting tumor mass. Since Flk-1 is believed to be

essential for formation of blood vessels, blocking Flk-1 activity by expression of a truncated receptor, might function to inhibit vascularization of the developing tumor and, thereby, inhibit its growth. As illustrated in Figures 13 and 14, coinjection of virus producing cells, expressing a truncated Flk-1 receptor, significantly inhibits the growth of the tumor as compared to controls receiving only tumor cells.

5.5. USE OF F1k-1 RECEPTOR OR LIGANDS

Receptor/ligand interaction between Flk-l and VEGF is believed to play an important role in the signalling system during vascularization and angiogenesis. Abnormal proliferation of blood vessels is an important component of a number of diseases.

Expression of Flk-1 RNA correlates with the development of the brain and with endothelial cell proliferation suggesting that Flk-1 might be a receptor involved in mediation of signaling events in the vascularization process. VEGF has been shown to be a mitogenic growth factor known to act exclusively on endothelial cell (Ferrara, N. and Henzel, W.J., 1989, Biochem. Biophys. Res. Comm. 161:851-858). Cross-linking and ligand binding experiments were performed, as described in Section 6.1.9 and 6.1.10 respectively, to determine whether VEGF is a ligand for Flk-1 and the results indicate that Flk-1 is an authentic high affinity

VEGF receptor (FIG 9).

In one embodiment of the invention, ligands for

Flk-1, the Flk-1 receptor itself, or a fragment
containing its VEGF binding site, could be administered
in vivo to modulate angiogenesis and/or vasculogenesis.
For example, administration of the Flk-1 receptor or a
fragment containing the VEGF binding site, could

competitively bind to VEGF and inhibit its interaction

with the native Flk-1 receptor in vivo to inhibit angiogenesis and/or vasculogenesis. Alternatively, ligands for Flk-1, including anti-Flk-1 antibodies or fragments thereof, may be used to modulate angiogenesis and/or vasculogenesis. Agonists of VEGF activity may be used to promote wound healing whereas antagonists of VEGF activity may be used to inhibit tumor growth.

Depending on the specific conditions being treated, these agents may be formulated and administered 10 systemically or locally. Techniques for formulation and administration may be found in "Remington's Pharmaceutical Sciences," Mack Publishing Co., Easton, PA, latest edition. Suitable routes may include oral, rectal, transmucosal, or intestinal administration; 15 parenteral delivery, including intramuscular, subcutaneous, intramedullary injections, as well as intrathecal, direct intraventricular, intravenous, intraperitoneal, intranasal, or intraocular injections, just to name a few. For injection, the agents of the 20 invention may be formulated in aqueous solutions, preferably in physiologically compatible buffers such as Hanks's solution, Ringer's solution, or physiological saline buffer. For such transmucosal administration, penetrants appropriate to the barrier to be permeated are 25 used in the formulation. Such penetrants are generally known in the art.

6. EXAMPLE: CLONING AND EXPRESSION PATTERNS OF F1k-1, A HIGH AFFINITY RECEPTOR FOR VEGF

The subsection below describes the cloning and characterization of the Flk-1 cDNA clone. Northern blot and in situ hybridization analyses indicate that Flk-1 is expressed in endothelial cells. Cross-linking and ligand binding experiments further indicate that Flk-1 is a high affinity receptor for VEGF.

6.1. MATERIALS AND METHODS

6.1.1. CDNA CLONING OF Flk-1

DNA extracted from \(\lambda\)gt10 cDNA library of day 8.5

mouse embryos (Fahrner et al., 1987, EMBO. J. 6:14971508) was used as template for polymerase chain reaction
(PCR; Saiki, R.K. et al., 1985 Science 230:1350-1354).

In an independent approach cDNA of capillary endothelial
cells that had been isolated from the brain of postnatal
day 4-8 mice was used for amplification (Risau, W., 1990)
In: development of the Vascular System. Issues Biomed.
Basel Karger 58-68 and Schnürch et al., unpublished)
Degenerated primers were designed on the basis of high
amino acid homologies within the kinase domain shared by
all RTKs (Wilks, A.F., 1989, Proc. Natl. Acad. Sci.
U.S.A. 86:1603-1607).

Full length cDNA clones of Flk-1 were isolated from another day 8.5 mouse embryo cDNA library, which had been prepared according to the method of Okayama and Berg (1983), and a day 11.5 mouse embryo λgt11 library (Clonetech) using the ³²P-labeled (Feinberg, A.P. and Vogelstein, B. 1983 Anal. Biochem. 132:6-13) 210-bp PCR fragment.

25 6.1.2. MOUSE EMBRYOS

Balb/c mice were mated overnight and the morning of vaginal plug detection was defined as 1/2 day of gestation. For Northern blot analysis the frozen embryos were homogenized in 5 M guanidinium thiocyanate and RNA was isolated as described (Ullrich, A. et al., 1985, Nature 313:756-761). For in situ hybridization, the embryos were embedded in Tissue-Tek (Miles), frozen on the surface of liquid nitrogen and stored at -70C prior to use.

6.1.3. PREPARATION OF PROBES

The 5'-located 2619 bp of the receptor cDNA were subcloned in the pGem3Z vector (Promega) as an EcoR1/ BamH1 fragment. The probe for Northern blot hybrid-5 ization was prepared by labelling the cDNA fragment with α -17PdATP (Amersham) by random hexanucleotide priming (Boehringer; Feinberg, A.P. and Vogelstein, B., 1983 Anal. Biochem. 132:6-13).

For in situ hybridization a single-strand antisense 10 DNA probe was prepared as described by Schnürch and Risau (Development, 1991 111:1143-54). The plasmid was linearized at the 3' end of the cDNA and a sense transcript was synthesized using SP6 RNA polymerase (Boehringer). The DNA was degraded using DNAase (RNAase 15 free preparation, Boehringer Mannheim). With the transcript, a random-primed cDNA synthesis with a α -35 dATP (Amersham) was performed by reverse transcription with MMLV reverse transcriptase (BRL). To obtain small cDNA fragments of about 100 bp in average suitable for in situ 20 hybridization, a high excess of primer was used. Subsequently the RNA transcript was partially hydrolyzed in 100 mM NaOH for 20 minutes at 70°C, and the probe was neutralized with the same amount of HC1 and purified with a Sephadex C50 column. After ethanol precipitation the 25 probe was dissolved at a final specific activity of 5x10' cpm. For control hybridization a sense probe was prepared with the same method.

RNA EXTRACTION AND NORTHERN ANALYSIS

Total cytoplasmic RNA was isolated according to the acidic phenol-method of Chromczynski and Sacchi (1987). $Poly(A^+)$ RNA aliquots were electrophoresed in 1.2% agarose formaldehyde (Sambrook, J. et al., 1989 Molecular Cloning: A Laboratory Manual 2nd ed. Cold Spring Harbor 35 Laboratory Press) gels and transferred to nitrocellulose

membranes (Schleicher & Schuell), Hybridizations were performed overnight in 50% formamide, 5 x SSC (750mM sodium chloride, 75mM sodium citrate), 5 x Denhardt's (0.1% Ficoll 400, 0.1% polyvinylpryollidone, 0.1% BSA) and -0.5% SDS at 42°C with 1-3x106 cpm-ml-1 of ^{NP}-Random primed DNA probe, followed by high stringency washes in 0.2 x SSC, 0.5% SDS at 52°C. The filters were exposed for 4 to 8 days.

6.1.5. IN SITU HYBRIDIZATION

Subcloning postfixation and hybridization was essentially performed according to Hogan et al. (1986). 10 μm thick sections were cut at -18°C on a Leitz cryostat. For prehybridization treatment no incubation with 0.2M HCl for removing the basic proteins was performed. Sections were incubated with the ³⁵S-cDNA probe (5x10⁴cpm/μ1) at 52°C in a buffer containing 50% formamide, 300 mM NuCl, 10 mM Tris-HCl, 10 mM NaPO4 (pH 6.8), 5 mM EDTA, 0.02% Ficoll 400, 0.01% polyvinylprolidone 0.02% BSA 10 m /ml yeast RNA, 10% dextran sulfate, and 10 mM NaCl, 10 mM Tris-HCl, 10 mM NaPO4 (pH 6.8), 5 mM EDTA, 10 mM DTT at 52°C). For

emulsion and exposed for eight days. After developing,

the sections were counterstained and toluidine blue or

May-Grinwald.

6.1.6. PREPARATION OF ANTISERA

autoradiography, slides were coated with Kodak NTB2 film

The 3' primed EcoRV/HindII fragment comprising the

128 C-terminal amino acids of Flk-1 was subcloned in the
fusion protein expression vector pGEX3X (Smith, D.B. and
Johnson, K.S., 1990 Gene. 67:31-40; Pharmacia). The
fusion protein was purified as described and used for
immunizing rabbits. After the second boost the rabbits

were bled and the antiserum was used for immunoprecipitation.

6.1.7. TRANSIENT EXPRESSION OF Flk-1 IN COS-1 CELLS

Transfection of COS-1 cells was performed essentially as described by Chen and Okayama (1987 Mol. Cell. Biol. 7:2745-2752) and Gorman et al. (1989 Virology 171:377-385). Briefly, cells were seeded to a density of 10 1.0 \times 106 per 10-cm dish and incubated overnight in DMEM containing 10% fetal calf serum (Gibco). 20 μ g of receptor cDNA cloned into a cytomegalovirus promotor driven expression vector was mixed in 0.5 ml of 0.25 M $CaCa_2$, 0.5 ml of 2 x BBS (280 mM NaCl, 1.5 mM Na_2HPO_4 , 50 15 mM BES, pH 6.96 and incubated for 30 min at room temperature. The calcium phosphate/DNA solution was then added to the cells, swirled gently, and incubated for 18 hours at 37°C under 3% CO2. For ligand binding experiments, the cells were removed from the plate and 20 treated as described below.

To obtain VEGF conditioned media, cells were transfected in 15-cm dishes. Media was collected after 48 h and VEGF was partially purified by affinity chromatography using heparin High Trap TM columns 25 (Pharmacia) and concentrated by ultrafiltration (Ferrara, N. and Henzel, W.J. 1989 Biochem. Biophys. Res. Comm. 161:851-858). The concentration of VEGF was determined by a ligand competition assay with bovine aortic endothelial cells.

For autophosphorylation assays, cells were seeded in 30 6-well dishes (2x10' cells per well), transfected as described above, and starved for 24 h in DMEM containing 0.5% fetal calf serum. The cells were then treated with 500 pM VEGF for 10 min. at 37°C or left untreated and 35 were subsequently lysed as described by Kris et al.

(1985). Flk-1 was immunoprecipitated with an antiserum raised in rabbits against the C-terminus of the receptor. The immunoprecipitates were separated on a 7.5% SDS polyacrylamide gel, transferred to nitrocellulose, and incubated with a mouse monoclonal antibody directed against phosphotyrosine (5E2; Fendly, B.M. et al., 1990 Cancer Research 50:1550-1558). Protein bands were visualized using horseradish peroxidase coupled goat anti-mouse antibody and the ECLTM (Amersham) detection system.

6.1.8. RADIOIODINATION OF VEGF

Recombinant human VEGF (5 μ g; generously provided by Dr. H. Weich) was dissolved in 110 μ l sodium phosphate 15 buffer pH 76, and iodinated by the procedure of Hunter and Greenwood (1962). The reaction products were separated from the labeled protein by passage over a sephadex G50 column, pre-equilibrated with phosphate buffered saline (PBS) containing 0.7% bovine serum 20 albumin (BSA), and aliquots of the collected fractions were counted before and after precipitation with 20% trichloracetic acid. The purity of the iodinated product was estimated to be superior to 90%, as determined by gel electrophoresis, and the specific activity was 77000 25 cpm/ng. The bioactivity of the iodinated VEGF was confirmed by comparison with the bioactivities of native VEGF using the tissue factor introduction assay described by Clauss, M. et al. (1990 J. Exp. Med. 172:1535-1545).

COS-1 cells transiently expressing Flk-1 and untransfected COS-1 cells were incubated with 200 pm 125I-VEGF at 4°C overnight, then washed twice with PBS and exposed to 0.5 mM disuccinimidyl suberate (DSS) in PBS

35 for 1 h at 4°C. The cells were lysed, Flk-1

immunoprecipitated, and analyzed by electrophoresis on a 7% polytarcylamide gel followed by autoradiography.

6.1.10. VEGF BINDING

Ligand binding experiments were performed as 5 described previously (Schumacher, R. et al., 1991, J. Biol. Chem. 266:19288-19295), COS-1 cells were grown in a 15-cm culture dish in DMEM for 48h after transfection. Cells were then washed carefully with PBS and incubated 10 with 5 ml of 25 mM EDTA in PBS for 10 min. Cells were then removed from the plate, washed once with binding buffer (DMEM, 25 mM HEPES, pH 7.5, 0.15% gelatin) and resuspended in 5 ml of binding buffer to determine the cell number. In a total volume of 500 μ l this cell 15 suspension was incubated for 90 min at 15°C with 10 pM 125 I-VEGF, and increasing concentration of unlabeled ligand (from 0 to 7 \times 10 9), which was partially purified from conditioned media of COS-1 cells transiently expressing VEGF (164 amino acid form; Breier et al., 1992). After 20 incubation, cells were washed with PBS 0.1% PBS in the cold. Free ligand was removed by repeated centrifugation and resuspension in binding buffer. Finally, the 125I radioactivity bound to the cells were determined in a gamma counter (Riastar). Data obtained were analyzed by 25 the method of Munson, P.J. and Rodbard, D. (1980 Anal. Biochem. 107:220-235).

6.1.11. RETROVIRAL VECTORS ENCODING TRANSDOMINANT-NEGATIVE MUTANTS OF Flk-1

Recombinant retroviral vectors were constructed that contained the coding region for amino acids 1 through 806 of the Flk-1 receptor (pLX Flk-1 cl.1 and cl.3, Figure 12). A recombinant virus containing a truncated c-fms receptor mutant (pNTK cfms TM cl.7) was used as a control. To obtain virus producing cells mouse GPE cells

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were infected with amphotrophic virus-containing conditioned media of PA317 cells that had been transfected with recombinant retroviral DNA. C6 gliobastoma tumor cells were implanted into nude mice either alone or coimplanted with virus producing cells. Injected cell numbers for the two sets of experiments are indicated below. Beginning at the time when the first tumors appeared, tumor volumes were measured every 2 to 3 days to obtain a growth curve.

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Experiment No. 1

Number of Mice	Number of C6 Cells	Virus-Producer Cell Line	Number of Virus-Cells
4	5 x 10 ³	pLXSN Flk-1 TM cl.3	1 x 10 ⁷
Δ	5 x 10 ⁵	None	0
4	5 x 10 ⁵	pNTK cfms TM cl.7	5 x 10 ⁶

Experiment No. 2

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4.	u

Number of	Number of C6 Cells	Virus-Producer Cell Line	Number of Virus-Cells
4	2 x 10°	pLXSN Flk-1 TM cl.1	2 x 10 ⁷
4	2 x 10°	pLXSN Flk-1 TM cl.3	2 x 10 ⁷
	2 x 10 ⁴	None	0
<u> </u>	2 x 10 ⁶	DNTK cfms TM cl.7	2 x 10 ⁷

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6.2. RESULTS

6.2.1. ISOLATION OF F1k-1

To identify RTKs that are expressed during mouse

development, PCR assays using two degenerate
oligonucleotide primer pools that were designed on the
basis of highly conserved sequences within the kinase
domain of RTKs were performed (Hanks, S.K. et al. 1988,
Science 241:42-52). DNA extracted from a \(\lambda\gamma\) CDNA
library of day 8.5 mouse embryos (Fahrner, K. et al.,

1987, EMBO. J., 6:1497-1508), a stage in mouse development at which many differentiation processes begin was used as the template in the PCR assays. In a parallel approach, with the intention of identifying RTKs 5 that regulate angiogenesis, similar primers were used for the amplification of RTK cDNA sequences from capillary endothelial cells that had been isolated from the brains of postnatal day 4-8 mice, a time at which brain endothelial cell proliferation is maximal (Robertson, 10 P.L. et al., 1985, Devel. Brain Res. 23:219-223). Both approaches yielded cDNA sequences (FIG. 11, SEQ. ID NO.:) encoding the recently described fetal liver RTK, Flk-1 (Matthews, W. et al., 1991, Proc. Natl. Acad. Sci. U.S.A. 88:9026-9030). Based on amino acid homology, this 15 receptor is a member of the type III subclass of RTKs (Ullrich, A. and Schlessinger, J. 1990, Cell 61:203-212) and is closely related to human flt, which also contains seven immunoglobin-like repeats in its extracellular domain in contrast to other RTKs of that subfamily, which 20 contain only five such repeat structures (Matthews, W. et al., 1991, Proc. Natl. Acad Sci. U.S.A. 88:9026-9030). Sequence comparisons of Flk-1 with KDR (Terman, B.I. et al., 1991, Oncogene 6:1677-1683) and TKr-C (Sarzani, R. et al., 1992, Biochem. Biophys. Res. Comm. 186:706-714) 25 suggest that these are the human and rat homologues of Flk-1, respectively (Figure 1).

6.2.2 EXPRESSION OF Flk-1 mRNA DURING EMBRYONIC DEVELOPMENT

As a first step towards the elucidation of the biological function of Flk-1, the expression of Flk-1 mRNA was analyzed in mouse embryos at different development stages. Northern blot hybridization experiments indicated abundant expression of a major 5.5 kb mRNA between day 9.5 and day 18.5, with an apparent

decline towards the end of gestation (Figure 2A). In postnatal day 4-8 brain capillaries Flk-1 mRNA was found to be highly enriched compared to total brain mRNA (Figure 2B).

- In situ hybridization experiments were performed to obtain more detailed information about the expression of Flk-1 during different embryonal stages. A singlestranded antisense, 2619-nucleotide-long DNA probe comprising the Flk-1 extracellular domain was used as a 10 probe because it generated the most specific hybridization signals. As an example, a parasagittal section of a day 14.5 embryo is shown in Figure 3. High levels of hybridization were detected in the ventricle of the heart, the lung, and the meninges; other tissues such 15 as brain, liver, and mandible appeared to contain fewer cells expressing Flk-1 mRNA. Thin strands of Flk-1 expression were also observed in the intersegmental regions of the vertebrae and at the inner surface of the atrium and the aorta. Higher magnification revealed that 20 the expression of Flk-1 seemed to be restricted to capillaries and blood vessels. Closer examination of the heart, for example, showed positive signals only in the ventricular capillaries and endothelial lining of the atrium (Figure 4A). In the lung, Flk-1 expression was 25 detected in peribronchial capillaries, but was absent from bronchial epithelium (Figure 4D). The aorta showed strong hybridization in endothelial cells, but not in the muscular layer (Figure 4C).
- The neuroectoderm in the telencephalon of a day 11.5 mouse embryo is largely avascular; the first vascular sprouts begin to radially invade the organ originating from the perineural vascular plexus (Bär, J., 1980, Adv. Anat. Embryol. Cell. Biol. 59:1-62; Risau, W. and Lemmon,

V. 1988, Dev. Biol. 125:441-450). At this stage, expression of Flk-1 was high in the perineural vascular plexus and in invading vascular sprouts, as shown in Figure 5A. These in situ hybridization analyses indicated that the 5 proliferating endothelial cells of an angiogenic sprout expressed the Flk-1 mRNA. At day 14.5, when the neuroectoderm is already highly vascularized, numerous radial vessels as well as branching vessels of the intraneural plexus contained large amounts of Flk-1 mRNA 10 (Figure 5B). At postnatal day 4, when sprouting and endothelial cell proliferation is at its highest, strong expression of Flk-1 mRNA was observed in endothelial cells (Figure 5C). Conversely, in the adult brain when angiogenesis has ceased, Flk-1 expression was very low 15 (Figure 5D) and appeared to be restricted mainly to the ehoroid plexus (Figure 6). In the choroid plexus, cells in the inner vascular layer expressed Flk-1 mRNA, while epithelial cells did not (Figure 6A, B).

The embryonic kidney is vascularized by an
angiogenic process (Ekblom, P. et al., 1982, Cell Diff.
11:35-39). Glomerular and peritubular capillaries
develop synchronously with epithelial morphogenesis. In
the postnatal day 4 kidney, in addition to other
capillaries, prominent expression of F1k-1 was observed
in the presumptive glomerular capillaries (Figure 7A).
This expression persisted in the adult kidney (Figure 7C
and D) and then seemed to be more confined to the
glomerular compared to the early postnatal kidney.

30 6.2.4. Flk-1 EXPRESSION IN ENDOTHELIAL CELL PROGENITORS

To investigate the possible involvement of Flk-1 in the early stages of vascular development, analysis of embryos at different stages during blood island formation 35 were performed. In a sagittal section of the deciduum of a day 8.5 mouse embryo, Flk-1 expression was detected on maternal blood vessels in the deciduum, in the yolk sac and in the trophectoderm. Flk-1 mRNA was also found in the allantois and inside the embryo, mainly located in 5 that part where mesenchyma is found (Figure 8A). At a higher magnification of the maternal deciduum, high levels of Flk-1 mRNA expression were found in the inner lining of blood vessels, which consist of endothelial cells (Figure 8B). In the yolk sac, hybridization signals were confined to the mesodermal layer, in which the hemangioblasts differentiate (Figure 8C). Figure 8D shows a blood island at higher magnification, in which the peripheral angioblasts expressed a high level of Flk-1 mRNA.

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6.2.5. Flk-1 IS A HIGH AFFINITY RECEPTOR FOR VEGF Detailed examination of in situ hybridization results and comparison with those for VEGF recently reported by Breier, G. et al. (1992, Development 114:521-20 532) revealed a remarkable similarity in expression pattern. Furthermore, Flk-1 expression in the glomerular endothelium and VEGF in the surrounding epithelial cells (Breier, G. et al., 1992, Development 114:521-532) raised the possibility of a paracrine relationship between these 25 cells types and suggested therefore a ligand-receptor relationship for VEGF and Flk-1, respectively. In order to test this hypothesis, the full-length Flk-1 cDNA was cloned into the mammalian expression vector pCMV, which contains transcriptional control elements of the human 30 cytomegalovirus (Gorman, C.M. et al., 1989, Virology 171:377-385). For transient expression of the receptor, the Flk-1 expressing plasmid was then transfected into COS-1 fibroblasts.

Specific binding of VEGF to the Flk-1 RTK was

35 demonstrated by crosslinking and competition binding

experiments. Purified ¹²⁵I-labeled VEGF was incubated with COS-1 cells transfected with the pCMV-Flk-1 expression vector. Crosslinking with DSS and subsequent analysis of immunoprecipitation, PAGE, and autoradiography revealed

- an approximately 220 kD band which was not detected in the control experiment with untransfected COS-1 cells and is likely to represent the VEGF/Flk-1 receptor complex (Figure 9A). In addition, VEGF competed with 125I-VEGF binding to Flk-1 expressing COS-1 cells (Figure 9B),
- whereas untransfected COS-1 cells did not bind ¹²⁵I-VEGF.

 The interaction of VEGF with the receptor on transfected cells was specific, as PDGF-BB did not compete with binding of ¹²⁵I-VEGF. Analysis of the binding data revealed a Kd of about 10⁻¹⁰ M, suggesting that Flk-1 is a
- high affinity receptor of VEGF. This finding, together with the Flk-1 and VEGF in situ hybridization results strongly suggests that Flk-1 is a physiologically relevantly receptor for VEGF.

An autophosphorylation assay was performed to
confirm the biological relevance of VEGF binding to the
Flk-1 receptor. COS1 cells which transiently expressed
Flk-1 were starved in DMEM containing 0.5% fetal calf
serum for 24h, stimulated with 0.5 mM VEGF, and lysed.
The receptors were immunoprecipitated with the Flk-1

- specific polyclonal antibody CT128, and then analyzed by SDS-PAGE and subsequent immunoblotting using the antiphosphotyrosine antibody 5E2 (Fendly, B.M. et al., 1990, Cancer Research 50:1550-1558). A shown in Figure 10, VEGF stimulation of Flk-1 expressing cells led to a
- 30 significant induction of tyrosine phosphorylation of the 180 kD Flk-1 receptor.

6.2.6. INHIBITION OF TUMOR GROWTH BY TRANSDOMINANT-NEGATIVE INHIBITION OF F1k-1

The Flk-1 receptor is believed to play a major role in vasculogenesis and angiogenesis. Therefore,

- 5 inhibition of Flk-1 activity may inhibit vasculogenesis of a developing tumor and inhibit its growth. To test this hypothesis, tumor cells (C6 rat glioblastoma) and mouse cells producing a recombinant retrovirus encoding a truncated Flk-1 receptor were mixed and implanted
- subcutaneously into nude mice. The implanted C6 glioblastoma cells secrete VEGF which will bind to and activate the Flk-1 receptors expressed on the surface of mouse endothelial cells. In the absence of any inhibitors of vasculogenesis, the endothelial cells will
- proliferate and migrate towards the tumor cells.

 Alternatively, if at the time of injection, the tumor cells are co-injected with cells producing recombinant retrovirus encoding the dominant-negative Flk-1, the endothelial cells growing towards the implanted tumor
- cells will become infected with recombinant retrovirus which may result in dominant-negative Flk-1 mutant expression and inhibition of endogenous Flk-1 signaling. Suppression of endothelial cell proliferation and migration will result in failure of the implanted tumor
- cells to become vascularized which will lead to inhibition of tumor growth. As shown in Figures 12 and 13, tumor growth is significantly inhibited in mice receiving implantations of cells producing truncated Flk-1 indicating that expression of a truncated Flk-1
- receptor can act in a dominant-negative manner to inhibit the activity of endogenous wild-type Flk-1.

The present invention is not to be limited in scope by the exemplified embodiments which are intended as illustrations of single aspects of the invention, and any clones, DNA or amino acid sequences which are

functionally equivalent are within the scope of the invention. Indeed, various modifications of the invention in addition to those described herein will become apparent to those skilled in the art from the foregoing description and accompanying drawings. Such modifications are intended to fall within the scope of the appended claims.

It is also to be understood that all base pair sizes given for nucleotides are approximate and are used for purposes of description.

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SEQUENCE LISTING

- (i) APPLICANT: Ullrich, et al
- (ii) TITLE OF INVENTION: FIX-1 IS A RECEPTOR FOR VASCULAR ENDOTHELIAL GROWTH FACTOR
- (iii) NUMBER OF SEQUENCES: 2
- (iv) CORRESPONDENCE ADDRESS:
 - (A) ADDRESSEE: Pennie & Edmonds
 - (B) STREET: 1155 Avenue of the Americas
 - (C) CITY: New York
 - (D) STATE: New York (E) COUNTRY: U.S.A.

 - (F) ZIP: 10036-2711
 - (V) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/HS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.25
- (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER: To be assigned
 - (B) FILING DATE: 03-MAR-1993 (C) CLASSIFICATION:
- (Viii) ATTORNEY/AGENT INFORMATION:
 - (A) NAME: Coruzzi, Laura A.
 - (B) REGISTRATION NUMBER: 30,742
 - (C) REFERENCE/DOCKET NUMBER: 7683-034-999
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 - (C) TELEX: 66141 PENNIE
- (2) INFORMATION FOR SEQ ID NO:1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 5470 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: unknown
 - (ii) MOLECULE TYPE: DNA (genomic)
 - (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION: 286..4386
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:
- TATAGGGCGA ATTGGGTACG GGACCCCCCT CGAGGTCGAC GGTATCGATA AGCTTGATAT 60 CGAATTCGGG CCCAGACTGT GTCCCGCAGC CGGGATAACC TGGCTGACCC GATTCCGCGG 120 ACACCGCTGA CAGCCGCGC TGGAGCCAGG GCGCCGGTGC CCCGGGTCTT 180 GCGCTGCGGG GGCCATACCG CCTCTGTGAC TTCTTTGCGG GCCAGGGACG GAGAAGGAGT 240

CTGI	CCCI	GA G	AAAC	TGGG	ic to	TGTG	CCCA	GGC	GCGA	GGT	GCAG	Мe		G AG u Se		294
AAG Lys	GCG Ala 5	CTG Leu	CTA Leu	GCT Ala	GTC Val	GCT Ala 10	CTG Leu	TGG Trp	TTC Phe	TGC Cys	GTG Val 15	GAG Glu	ACC Thr	CGÀ Arg	GCC Ala	342
GCC Ala 20	TCT Ser	GTG Val	GCT Gly	TTG Leu	ACT Thr 25	ely eec	GAT Asp	TTT Phe	CTC Leu	CAT His 30	CCC Pro	CCC Pro	AAG Lys	CTC Leu	AGC Ser 35	390
ACA Thr	CAG Gln	AAA Lys	yab	ATA Ile 40	CTG Leu	ACA Thr	ATT Ile	TTG Leu	GCA Ala 45	TAA nbA	ACA Thr	ACC Thr	CTT Leu	CAG Gln 50	ATT Ile	438
ACT Thr	TGC Cys	AGG Arg	GGA Gly 55	CAG Gln	CGG Arg	GAC Asp	CTG Leu	GAC Asp 60	TGG Trp	CTT Leu	TGG Trp	CCC Pro	AAT Asn 65	GCT Ala	CAG Gln	486
CGT Arg	GAT Asp	TCT Ser 70	GAG Glu	GAA Glu	AGG Arg	GTA Val	TTG Leu 75	GTG Val	ACT Thr	GAA Glu	TGC Cys	GGC Gly 80	GGT Gly	GGT Gly	yab GyC	S34
AGT Ser	ATC :Ile 85	TTC Phe	TGC Cys	AAA Lys	ACA Thr	CTC Leu 90	ACC Thr	ATT Ile	CCC Pro	AGG Arg	GTG Val 95	GTT Val	GGA Gly	AAT Asn	GAT Asp	582
ACT Thr 100	Gly	GCC Ala	TAC Tyr	AAG Lys	TGC Cys 105	TCG Ser	TAC Tyr	CGG Arg	GAC Asp	GTC Val 110	GAC Asp	ATA Ile	GCC Ala	TCC Ser	ACT Thr 115	630
CTT Val	TAT Tyr	GTC Val	TAT	GTT Val 120	CGA	GAT Asp	TAC	AGA Arg	TCA Ser 125	CCA Pro	TTC Phe	ATC Ile	GCC Ala	TCT Ser 130	GTC Val	678
AGT Ser	Asp	CAG Gln	CAT His 135	Gly	ATC Ile	GTG Val	TAC	ATC Ile 140	ACC Thr	GAG Glu	AAC Asn	AAG Lys	AAC Asn 145	AAA Lys	ACT Thr	726
GTG Val	GTG Val	ATC Ile 150	Pro	TGC Cys	CGA Arg	gjå ecc	TCG Ser 155	Ile	TCA Ser	AAC Asn	CTC Leu	AAT Asn 160	Val	TCT Ser	CTT	774
Cys	GCT Ala 165	Arg	TAT	CCA Pro	GAA Glu	AAG Lys 170	Arg	TTT Phe	GTT Val	CCG Pro	GAT Asp 175	Cly	AAC Asn	AGA Arg	ATT	822
TCC Ser 180	Trr	Asp GAC	AGC Ser	GAG Glu	ATA Ile 185	Gly	TTT Phe	ACT Thr	CTC	Pro 190	Ser	TAC	ATG Met	ATC	Ser 195	870
TAT	GCC Ala	Gly	ATG Met	GTC Val 200	. Phe	TGT Cys	GAG Glu	GCA Ala	AAG Lys 205	Ile	AAT Ast	GAT Asp	GAA Glu	Thr 210	TAT Tyr	918
CAC Glr	TCI Ser	ATC Ile	ATC Met 215	Tyr	ATA	GTI Val	GTC Val	GTT Val 220	. Val	GGA Gly	TAT Tyr	AGG Arg	11e 229	Ty	GAT Asp	966
GT(Va)	ATT	CTC Let 230	ı Ser	CCC Pro	C CCC	CAT His	GAJ G10 23	ı Ile	GAC Glu	CTA Lev	TC1	GCC Ala 240	Gly	GAI Glu	TA8	1014
CT: Le	CTC	l Le	CAA A	TG:	s Thi	r Ala	a Ar	A ACI	: Glu	ı Lei	ı Ası	ya.	G GGG	CT:	I GAT	106

								-5	2-								
TTC Phe 260	ACC Thr	TCG Trp	HTR CYC	TCT Ser	CCA Pro 265	CCT Pro	TCA Ser	aag Lys	Ser	CAT His 270	CAT His	aag Lys	aag Lys	ATT Ile	GTA Val 275	•	1110
AAC ABO	CGG Arg	CAI Asp	GTG Val	AAA Lys 280	Pro CCC	TIT Phe	CCT Pro	CCC	ACT Thr 285	GTG Val	GCG Ala	AAG Lys	ATG Ket	TTT Phe 290	TIG	; 1	1158
AGC Ser	ACC Thr	Leu	ACA Thr 295	ATA Ile	GAA Glu	AGT Ser	GTG Val	ACC Thr 300	AAG Lys	AGT Ser	GAC Asp	CAA Gln	GGG Gly 305	GAA Glu	Typ		1206
ACC Thr	TGT Cys	GTA Val 310	Ala	TCC Ser	AGT Ser	GGA Gly	CGG Arg 315	ATG Ket	ATC Ile	AAG Lys	Arg	AAT Asn 320	, wed	ACA Thr	Phe	:	1254
GTC Val	CGA Arg 325	GTT Val	CAC	ACA Thr	AAG Lys	CCT Pro 330	TTT Phe	ATT Ile	GCT Ala	TTC Phe	GGT Gly 335	AGT Ser	GCG	ATG Met	AA! Lyi	A B	1302
TCT Ser 340	Leu	GTG Val	GAA Glu	GCC	ACA Thr 345	GTG Val	GGC	AGT Ser	CAA Gln	GTC Val 350	Arg	ATC	CCT Pro	GTG Val	Ly: 35	D	1350
TAT Tyr	CTC Leu	AGT	TAC	CCA Pro	GCT Ala	CCT	GAT Asp	ATC : Ile	AAA Lys 365	Trp	TAC	AGA Arg	AAT ABD	GGA Gly 370	M.E	g ·	1398
CCC	ATT	GAG Glu	TCC Ser 375	Asr	TAC Tyr	ACA Thr	ATG Het	Ile 380	Val	GGC	GAT Asp	GAF Glu	CTC Lev 385	Tur	AT Il	C .e	1446
ATG Met	GAA Glv	GT(Va) 39(Thi	GAZ Glu	A AGA 1 Arg	GAT Asp	GCA Ala 395	Gly	AAC Asn	TAC	ACC Thi	GT(r TT4	CTC	AC 1 Th	ic ir	1494
AAC Asn	CCC Pro 405	Ile	r TCI e Sei	A ATO	G GAG	AAF Lys 410	Glr	a Ser	CAC His	ATO Met	GTC E Val 419	r se	r Cro	G GT	r G1 l Va	r) C	1542
AA1 As: 420	va.	C CC	A CC	C CAC	G ATO	e Gly	r GAG	AAI Lyi	A GCC	1 Le	a 110	C TC e Se	c cc r Pr	T AT	L A	AT SP 35	1590
TC(Se:	TA:	C CA	G TA'	T GG T G1 44	G ACC Y Th	C ATO	G CAG	G ACI	A TTO Let 445	ı Th	A TG r Cy	C AC	A GT r Va	C TA 1 Ty 45	EA	CC la	1638
AA Asi	C CC	r cc o Pr	C CT o Le 45	u Hi	C CA s Hi	C AT	C CA	G TG n Tr 46	p Ty	C TG F Tr	g CA p Gl	G CI n Le	A GA nu Gl 46	u Gi	A G	cc la	1686
TG Cy	C TC B Se	C TA r Ty 47	r Ar	A CC	C GG	C CA y Gl	A AC n Th 47	r Se	c cc	G TA o Ty	r ģl	.a cy	T AP /S Ly 30	A GF	LU T	rp CC	1734
AG Ar	A CA g Hi 48	g Va	G GA	G GA .u As	T TI p Ph	C CA le Gl 49	n Gl	g gg y gl	A AA y As	C AA n Ly	G A1 /s I1 49	.e G.	AA G7 Lu Va	C AC	ec A	lAA Lys	1782
AA A a 50	n Gl	A TI	T GO	cc cr	IA DI LI DE SC	Le Gl	la Go	A AA Y Ly	AA AA BA B'	in L	AA AC /B Th	or G	TA AG	ST A	ur i	CTG Leu 515	. 1830
G1 Va	C AT	C C	AA GO ln Al	la A	CC AF la Ar 20	AC G1 an Va	IG TO	CA GO er Al	CG TI La Le 52	u T	AC AI Yr L	AA T ys C	Ya C	Lu n	CC 1 1a : 30	ATC Ile	1878

AAC Asn	AAA Lys	GCG Ala	GGA Gly 535	CGA Arg	GGA Gly	GAG Glu	AGG Arg	GTC Val 540	ATC Ile	TCC Ser	TTC Phe	CAT His	GTG Val 545	ATC Ile	AGG Arg	1926
					GTG Val											1974
					TGC Cys							Phe				2022
					GGC Gly 585											2070
TCA Ser	CTC Leu	ACA Thr	CCA Pro	GTT Val 600	TGC Cys	AAG Lys	AAC Asn	TTG Leu	GAT Asp 605	GCT Ala	CTT Leu	TGG Trp	AAA Lys	CTG Leu 610	AAT Asn	2118
					AAC Asn											2166
					CAG Gln										CAA .Gln	2214
					AAA Lys											2262
					CCC Pro 665											2310
															AAT Asn	2358
									Asp						GAA Glu	2406
			Ile					Gly					Thr		CGC	2454
		Arg					Gly					Gln			AAT Asn	2502
GTC Val 740	Leu	GLY	Cys	GCA Ala	AGA Arg 745	Ala	GAG Glu	ACG Thr	CTC Leu	Phe 750	Ile	ATA Ile	GAA Glu	GCT	GCC Ala 755	2550
CAG Gln	GAA Glu	AAG Lys	ACC	AAC Asn 760	Leu	GAA Glu	GTC Val	ATT Ile	TILE 765	Leu	GTC Val	GGC Gly	ACT Thr	GCA Ala 770	GTG Val	2598
				Phe					ı Val					Thi	GTT Val	2646
AAG Lys	CGG	GCC Ala 790	Asr	GAP Glu	GGG Gly	GAA Glu	CTC Lev 795	Lys	ACA Thi	GLY	TAC Tyr	TTC Lev 800	ı Ser	ATT	GTC Val	2694

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ATG Het	GAT Asp 805	CCA Pro	GAT Asp	GAA Glu	TTG Leu	CCC Pro 810	TTG Leu	CAT Asp	GAG Glu	CGC Arg	TGT Cys 815	GAA Glu	yrg	TTG Leu	CCT Pro	2742
TAT Tyr 820	GAT Asp	GCC Ala	AGC Ser	Lys	TGG Trp 825	GAA Glu	TTC Phe	CCC Pro	agg Arg	GAC Asp 830	CGG Arg	CTG Leu	AAA Lys	CTA Leu	GGA Gly 835	2790
YYY Ta	CCT Pro	CTT Leu	GGC Gly	CGC Arg 840	GGT Gly	GCC Ala	TIC Phe	ejå eec	CAA Gln 845	GTG Val	ATT Ile	GAG Glu	GCA Ala	GAC Asp 850	GCT Ala	2838
TTT Phe	GGA Gly	ATT Ile	GAC Asp 855	AAG Lys	ACA Thr	GCG Ala	ACT Thr	TGC Cys 860	AAA Lys	ACA Thr	GTA Val	gcc Ala	GTC Val 865	AAG Lys	ATG Met	2886
TTG Leu	AAA Lys	GAA Glu 870	Cly	GCA Ala	ACA Thr	CAC	AGC Ser 875	GAG Glu	CAT His	CGA Arg	GCC Ala	CTC Leu 880	Het	TCT Ser	GAA Glu	2934
CTC	AAG Lys 885	Ile	CTC	ATC	CAC His	ATT Ile 890	GGT	CAC His	CAT His	CTC Leu	AAT Asn 895	GTG Val	GTG Val	AAC	CTC	2982
CTA Leu 900	Gly	GCC	TGC Cys	ACC Thr	AAG Lys 905	Pro	GGA Gly	es eee	CCT	CTC Leu 910	wet	GTG Val	ATT	GTG Val	GAA Glu 915	3030
TTC	TGC Cys	AAG Lys	TTT	GGA Gly 920	Asn	CTA Leu	TCA Ser	ACT	TAC Tyr 925	Leu	CGG Arg	GC	AAG Lys	AGA Arg 930	AAT Asn	3078
GAA Glu	TTI Phe	GTI Val	935	Tyr	AAG Lys	AGC Ser	AAA Lys	GGG Gly 940	Ala	CGC Arg	TTC Phe	CGC Arg	CAG G G1: 949	GTZ	AAG Lya	3126
GA(TAC Tyl	GT1 Val	L Gly	GAG Glu	CTC	TCC Ser	GTG Val	Yei	CTC Lev	AAA Lys	AGF Arq	A CGG	a red	GA(C AGC p Ser	3174
ATC Ile	C ACC e Thi 96	r Se	C AGO	CAC CGl	AGC A Sei	TC1 Ser 970	Ala	Z AGO A Sei	C TCI r Se:	GGC Gly	7 Phe 97	e va	T GAG	G GA	TAN TAN	3222
TC: Se: 98:	r Le	C AG	GA:	r GT	A GAG 1 G1: 98:	i Cli	GAI Gl	A GAI	A GC: u Ala	I IC	C GI	A GA u Gl	A CT	G TA u Ty	C AAG r Lys 995	3270
GA As	C TT p Ph	C CT e Le	G AC u Th	C TT r Le	u Gl	G CA:	CT Le	C AT u Il	e Cy	T TAG 8 Ty: 05	C AG	C TT r Ph	C CA e Gl	A GT n Va 10	G GCT 1 Ala 10	3318
AA Ly	G GG	C AT y He	t Gl	G TT u Ph 15	C TT e Le	G GC: u Al	a TC a Se	r Ar	G AA g Ly 20	G TG	T AT B Il	C CA e Hi	.s Ar	G GA g As	c CTG	3366
GC Al	A GC	a Ar	AA A g As OE	C AT	T CI e Le	C CI	u Se	G GA r Gl	G AA u Ly	G AA 8 As	T GI n Va	T AS	TT AR al Ly 040	G AT	C TGT Le Cys	3414
GA Ae	sp Pi	C GG ne G1	C TI Y Le	G GC	C CC	d ya	C A1	T II le Ty	T AF	A GA	ib Li	G G/ 10 A: 055	AT TA	NT G: /E V	C AGA al Arg	3462
L	AA GG YB G: 060	GA GA Ly Ad	AT GO	CC CC	g Le	C CC eu Pr 065	T TT	rg al Bu Ly	AG TO YS TI	cp we	rg go et Al 070	CC CC La P	CG GI	AA A	cc ATT hr Ile 107	

			Val		Thr					Val				1030 61Å 661	Val	3558
TTG Leu	CTC Leu	TCG Trp	GAA Glu 1095	Ile	TTT Phe	TCC Ser	TTA Leu	GGT Gly 1100	Ala	TCC Ser	CCA Pro	TAC Tyr	CCT Pro 1109	GCG GCG	GTC Val	3606
AAG Lye	ATT Ile	GAT Asp 1110	Çlu	GAA Glu	TTT Phe	TGT Cys	AGG Arg 1115	λrg	TTG Leu	AAA Lys	GAA Glu	GGA Gly 1120	The	AGA Arg	ATG Het	3654
CGG Arg	GCT Ala 1125	Pro	GAC Asp	TAC Tyr	ACT Thr	ACC Thr 1130	Pro	GAA Glu	ATG Het	TAC Tyr	CAG Gln 113	Thr	ATG Met	CTG Leu	GAC Asp	3702
TGC Cys 1140	Trp	CAT His	GAG Glu	GAC Asp	CCC Pro 1145	Asn	CAG Gln	AGA Arg	CCC Pro	TCG Ser 1150	Phe	TCA Ser	GAG Glu	TTG Leu	GTG Val 1155	3750
GAG Glu	CAT His	TTG Leu	GGA Gly	AAC Asn 1160	Leu	CTG Leu	CAA Gln	GCA Ala	AAT Asn 1169	Ala	CAG Gln	CAG Gln	GAT Asp	GGC Gly 1170	Lys	3798
GAC Asp	TAT Tyr	ATT Ile	GTT Val 117	Leu	CCA Pro	ATG Met	TCA Ser	GAG Glu 118	Thr	CTG	AGC Ser	ATG Met	GAA Glu 118	GAG Glu 5	GAT Asp	3846
TCI Ser	GGA Gly	CTC Leu 1190	Ser	CTG Leu	CCT Pro	ACC Thr	TCA Ser 119	Pro	GTT Val	TCC Ser	Cys	ATG Met 120	Glu	GAA Glu	GAG Glu	3894
GAA Glu	GTG Val 120	Сув	yab	CCC Pro	ДДД	TTC Phe 121	His	TAT Tyr	GAC Asp	AAC Asn	ACA Thr 121	Ala	GGA	ATC Ile	AGT Ser	3942
CAT His 122	Tyr	CTC Leu	CAG Gln	AAC Asn	AGT Ser 122	Lys	CGA Arg	AAG Lys	AGC Ser	ccc Arg 123	Pro	GTG Val	AGT Ser	GTA Val	AAA Lys 1235	3990
ACA Thr	TTT Phe	GAA Glu	GAT Asp	ATC Ile 124	Pro	TTG Leu	GAG Glu	GAA Glu	CCA Pro 124	Glu	GTA Val	AAA Lys	GTG Val	Ile 125	CCA Pro 0	4038
GAT Asp	GAC	AGC Ser	CAG Gln 125	Thr	GAC Asp	AGT Ser	GGG	ATG Met 126	Val	CTI Leu	GCA Ala	TCA Ser	GAA Glu 126	GIU	CTG Leu	4086
AAA Lys	ACT	CTG Leu 127	Glu	GAC	AGG Arg	AAC Asn	127	Lev	TCT Ser	CCA Pro	TCI Ser	TT1 Phe 128	GL	GGA Gly	ATG Het	4134
ATG Het	CCC Pro	Ser	AAA Lys	AGC Ser	AGG Arg	GAG Glu 129	Ser	Val	GCC L Ala	Ser	GA: Glv 129	ı Gly	TCC Sei	AAC Asr	CAG Gln	4182
ACC Thr 130	Ser	GLY	TAC Tyr	CAG Gln	TCI Ser 130	Gly	TAI	CAC His	TC? Sei	GA: A8; 13:) yei	ACI Thi	A GAG	D ACC	ACC Thr 1315	4230
GTC Val	TAC Tyr	TCC Ser	AGC Ser	GAC Asp 132	Glu	GC Ala	GG Gl	A. CT: y Let	r TTX Lev 13:	Ly:	G ATO	G GTG	G GA' l As	r GC: p Ala 13:	r GCA a Ala 30	4278
GT: Val	CAC His	C GCT	GAC Asp	Sei	A GGG	Th	Th:	r Le	G CAG	n Le	u Th	r Se	r Cy	s Le	TAA A	432

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GGA AGT GGT CCT GTC CCG GCT CCG CCC CCA ACT CCT GGA AAT CAC GAG Gly Ser Gly Pro Val Pro Ala Pro Pro Pro Thr Pro Gly Asn His Glu 1350 1355 1360	4374
AGA GGT GCT GCT TAGATTTTCA AGTGTTGTTC TTTCCACCAC CCGGAAGTAG Arg Gly Ala Ala 1365	4426
CCACATITGA TITTCATTIT TGGAGGAGGG ACCTCAGACT GCAAGGAGCT TGTCCTCAGG	4486
GCATTTCCAG AGAAGATGCC CATGACCCAA GAATGTGTTG ACTCTACTCT	4546
CATTTAAAAG TCCTATATAA TGTGCCCTGC TGTGGTCTCA CTACCAGTTA AAGCAAAAGA	4606
CTTTCAAACA CGTGGACTCT GTCCTCCAAG AAGTGGCAAC GGCACCTCTG TGAAACTGGA	4666
TCGAATGGGC AATGCTTTGT GTGTTGAGGA TGGGTGAGAT GTCCCAGGGC CGAGTCTGTC	4726
TACCTIGGAG GCTTTGTGGA GGATGCGGGC TATGAGCCAA GTGTTAAGTG TGGGATGTGG	4786
ACTGGGAGGA AGGAAGGCGC AAGTCGCTCG GAGAGCGGTT GGAGCCTGCA GATGCATTGT	4846
GCTGGCTCTG GTGGAGGTGG GCTTGTGGCC TGTCAGGAAA CGCAAAGGCG GCCGGCAGGG	4906
TITGGTTTTG GAAGGTTTGC GTGCTCTTCA CAGTCGGGTT ACAGGCGAGT TCCCTGTGGC	4966
GTTTCCTACT CCTAATGAGA GTTCCTTCCG GACTCTTACG TGTCTCCTGG CCTGGCCCCA	5026
GGAAGGAAAT GATGCAGCTT GCTCCTTCCT CATCTCTCAG GCTGTGCCTT AATTCAGAAC	5086
ACCARAGAG AGGAACGTCG GCAGAGGCTC CTGACGGGGC CGAAGAATTG TGAGAACAGA	5146
ACAGAAACTC AGGGTTTCTG CTGGGTGGAG ACCCACGTGG CGCCCTGGTG GCAGGTCTGA	5206
GGGTTCTCTG TCAAGTGGCG GTAAAGGCTC AGGCTGGTGT TCTTCCTCTA TCTCCACTCC	5266
TGTCAGGCCC CCAAGTCCTC AGTATTTTAG CTTTGTGGCT TCCTGATGGC AGAAAAATCT	5326
TARTIGGTIG GTTIGCTCTC CAGATAATCA CTAGCCAGAT TTCGAAATTA CTTTTTAGCC	5386
GAGGITATGA TAACATCTAC TGTATCCTTT AGAATTTTAA CCTATAAAAC TATGTCTACT	5446
GGTTTCTGCC TGTGTGCTTA TGTT	5470

(2) INFORMATION FOR SEQ ID NO:2:

- (i) SEQUENCE CHARACTERISTICS:
 (A) LENGTH: 1367 amino acids
 (B) TYPE: amino acid
 (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

Met Glu Ser Lys Ala Leu Leu Ala Val Ala Leu Trp Phe Cys Val Glu 1 5 15

Thr Arg Ala Ala Ser Val Gly Leu Thr Gly Asp Phe Leu His Pro Pro

Lys Leu Ser Thr Gln Lys Asp Ile Leu Thr Ile Leu Ala Asn Thr Thr

Leu Gln Ile Thr Cys Arg Gly Gln Arg Asp Leu Asp Trp Leu Trp Pro S0 S5 60

Asn Ala Gln Arg Asp Ser Glu Glu Arg Val Leu Val Thr Glu Cys Gly 65 70 75 80 Gly Gly Asp Ser Ile Phe Cys Lys Thr Leu Thr Ile Pro Arg Val Val 85 90 95 Gly Asn Asp Thr Gly Ala Tyr Lys Cys Ser Tyr Arg Asp Val Asp Ile 100 105 110 Ala Ser Thr Val Tyr Val Tyr Val Arg Asp Tyr Arg Ser Pro Phe Ile 115 120 125 Ala Ser Val Ser Asp Gln His Gly Ile Val Tyr Ile Thr Glu Asn Lys 130 140 Asn Lys Thr Val Val Ile Pro Cys Arg Gly Ser Ile Ser Asn Leu Asn 145 150 155 160 Val Ser Leu Cys Ala Arg Tyr Pro Glu Lys Arg Phe Val Pro Asp Gly 165 170 175 Asn Arg Ile Ser Trp Asp Ser Glu Ile Gly Phe Thr Leu Pro Ser Tyr Met Ile Ser Tyr Ala Gly Met Val Phe Cys Glu Ala Lys Ile Asn Asp 195 200 205 Glu Thr Tyr Gln Ser Ile Met Tyr Ile Val Val Val Gly Tyr Arg Ile Tyr Asp Val Ile Leu Ser Pro Pro His Glu Ile Glu Leu Ser Ala 225 230 235 240 Gly Glu Lys Leu Val Leu Asn Cys Thr Ala Arg Thr Glu Leu Asn Val 245 250 255 Gly Leu Asp Phe Thr Trp His Ser Pro Pro Ser Lys Ser His His Lys 260 265 270 Lys Ile Val Asn Arg Asp Val Lys Pro Phe Pro Gly Thr Val Ala Lys Met Phe Leu Ser Thr Leu Thr Ile Glu Ser Val Thr Lys Ser Asp Gln290295 Gly Glu Tyr Thr Cys Val Ala Ser Ser Gly Arg Ket Ile Lys Arg Asn 305 310 315 Arg Thr Phe Val Arg Val His Thr Lys Pro Phe Ile Ala Phe Gly Ser 325 330 335 Gly Met Lys Ser Leu Val Glu Ala Thr Val Gly Ser Gln Val Arg Ile Pro Val Lys Tyr Leu Ser Tyr Pro Ala Pro Asp Ile Lys Trp Tyr Arg 355 360 365 Asn Gly Arg Pro Ile Glu Ser Asn Tyr Thr Het Ile Val Gly Asp Glu Leu Thr Ile Met Glu Val Thr Glu Arg Asp Ala Gly Asn Tyr Thr Val 385 390 395 400 Ile Leu Thr Asn Pro Ile Ser Het Glu Lys Gln Ser His Het Val Ser Leu Val Val Asn Val Pro Pro Cln Ile Gly Glu Lys Ala Leu Ile Ser

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			420					425					430		
Pro	Ket	Авр 435	Ser	Tyr	Gln	Tyr	Gly 440	Thr	Xet	Gln	Thr	Leu 445	Thr	Сув	Thr
Val	Tyr 450	Ala	Asn	Pro	Pro	Leu 455	His	His	Ile	Gln	Trp 460	Tyr	Trp	Gln	Leu ·
Glu 465	Glu	Ala	Сув	Ser	Tyr 470	Arg	Pro	Gly	Gln	Thr 475	Ser	Pro	Tyr	Ala	Cys 480
				485	Val				490					495	
			500		Tyr			505					510		
		515			Gln		520					525			
	530				Ala	535					540				
545					Glu 550					555					560
				565	Ser				570					575	
			580		Tyr			585					590		
	_	595			Thr		600					605			
_	610					615					620				Ile
625					630					635					640
				645					650)				655	i
			660)				665	5				670)	Glu
		679	5	••••	•		680	1				689	5) Ala
	690)				695	•				700	,			1 Thr
709	5				710)				71	5				720
				729	5				73	0				/3:	
			74	0				74	5				75	Ų	e Ile
		75	5				760	0				76	5		l Gly
Th	E Al		1 11	e Al	a Met	2 Pho 77		e Tr	p Le	u Le	u Le 78	u Va O	1 11	e Va	l Leu

Arg 785	Thr	Val	Lys	Arg	Ala 790	Asn	Glu	Gly	Glu	Leu 795	Lys	Thr	Gly	Tyr	Leu 800
Ser	Ile	Val	Ket	Asp 805	Pro	Asp	Glu	Leu	Pro 810	Leu	Asp	Glu	Arg	Сув 815	Glu
Arg	Leu	Pro	Tyr 820	Yab	Ala	Ser	Lys	Trp 825	Glu	Phe	Pro	Arg	Asp 830	Arg	Leu
Lys	Leu	Gly 835	ŗĀŘ	Pro	Leu	Gly	Arg 840	Gly	Ala	Phe	Gly	Gln 845	Val	Ile	Glu
Ala	Asp 850	Ala	Phe	Gly	Ile	Asp 855	Lye	Thr	Ala-	Thr	Сув 860	Lys	Thr	Val	Ala
Val 865	Lys	Ket	Leu	Lys	Glu 870	Gly	Ala	Thr	His	Ser 875	Glu	His	Arg	Ala	Leu 880
Met	Ser	Glu	Leu	Lys 885	Ile	Leu	Ile	His	Ile 890	Gly	His	His	Leu	Asn 895	Val
Val	Asn	Leu	Lėu 900	Gly	Ala	Сув	Thr	Lys 905	Pro	Gly	Gly	Pro	Leu 910	Het	Val
Ile	Val	Glu 915	Phe	Сув	Lys	Phe	Gly 920	Asn	Leu	Ser	Thr	Tyr 925	Leu	Arg	Gly
Lys	Arg 930	Asn	Glu	Phe	Val	Pro 935	Tyr	Lys	Ser	Lys	Gly 940	λla	Arg	Phe	Arg
Gln 945	Gly	Lys	Asp	Tyr	Val 950	Gly	Glu	Leu	Ser	Val 955	Asp	Leu	Lys	Arg	960
Leu	Asp	Ser	Ile	Thr 965	Ser	Ser	Gln	Ser	Ser 970		Ser	Ser	Gly	Phe 975	Val
Glu	Glu	Lys	Ser 980	Lau	Ser	Asp	Val	Glu 985	Glu	Glu	Glu	Ala	Ser 990	Glu	Glu
Leu	Tyr	995		Phe	Leu	Thr	Leu 100		His	Leu	Ile	Cys 100	Tyr	Ser	Phe
Gln	Val 101		Lys	Cly	Ket	Glu 101		Leu	Ala	. Ser	Arg 102		Сув	Ile	His
Arg 102		Lev	Ala	Ala	Arg 103		Ile	Leu	Leu	Ser 103		Lys	Asn	Val	Val 1040
Lys	Ile	: Сув	Asp	Phe 104		Leu	Ala	Arg	105	Ile 0	Tyr	Lys	Asp	Pro 105	S S
Ty	. Val	Arg	106		ysb	Ala	Arg	106	Pro	Leu	Lys	Tr	107	Ala O	Pro
Glu	Thi	10		Asp	Arg	Val	108	Thi	: Ile	Glr	ı Ser	108	Va] 35	Tr	Ser
Phe	109		LLeu	Lev	Trp	109		Phe	e Ser	c Lev	110		a Sei	Pro	Tyr
Pro		y Va	L Lys	Ile	Asp 111		ı Glu	ı Phe	e Cys	11:	g Arq 15	g Le	ı Ly:	s Gl	1 Gly 1120
Th	r Ar	g Ke	t Arq	Ala 112	a Pro 25) As	р Туі	c Th	11:		o Glu	ı Me	ту:	r Gl:	n Thr 35
Me	t Le	u As	р Су	Tr:	e Hi	E Gl	u Asj	o Pr	o As	n Gl	n Ar	g Pr	o Se	r Ph	e Ser

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		1140					1145					1150				
Glu	Leu	Val 1155		His	Leu	Gly	Asn 1160		Leu	Gln	Ala	λsn 1165		Gln	Gln	
Asp				Tyr		Val 1175		Pro	Het	Ser	Glu 1180		Leu	Ser	Met	
Glu 1189		Дар	Ser	Gly	Leu 1190		Leu	Pro	Thr	Ser 1199		Val	Ser	Сув	Met 1200	
Glu	Glu	Glu	Glu	Val 1205		Asp	Pro	Lys	Phe 1210		Tyr	увр	Asn	Thr 1215		
Gly	Ile		His 1220	Tyr									Arg 1230		Val	

Ser Val Lys Thr Phe Glu Asp Ile Pro Leu Glu Glu Pro Glu Val Lys 1235 1240 1245

Val Ile Pro Asp Asp Ser Gln Thr Asp Ser Gly Met Val Leu Ala Ser 1250 1255 1260

Glu Glu Leu Lys Thr Leu Glu Asp Arg Asn Lys Leu Ser Pro Ser Phe 1265 1270 1275 1280

Gly Gly Met Met Pro Ser Lys Ser Arg Glu Ser Val Ala Ser Glu Gly 1285 1290 1295

Ser Asn Gln Thr Ser Gly Tyr Gln Ser Gly Tyr His Ser Asp Asp Thr 1300 1305 1310

Asp Thr Thr Val Tyr Ser Ser Asp Glu Ala Gly Leu Leu Lys Met Val

Asp Ala Ala Val His Ala Asp Ser Gly Thr Thr Leu Gln Leu Thr Ser 1330 1340

Cys Leu Asn Gly Ser Gly Pro Val Pro Ala Pro Pro Pro Thr Pro Gly 1345 1350 1355 1360

Asn His Glu Arg Gly Ala Ala 1365

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WHAT IS CLAIMED IS:

- A recombinant DNA vector containing a
 nucleotide sequence that encodes a Flk-1 operatively
 associated with a regulatory sequence that controls gene
 expression in a host.
- A recombinant DNA vector containing a nucleotide sequence that encodes a Flk-1 fusion protein
 operatively associated with a regulatory sequence that controls gene expression in a host.
 - 3. An engineered host cell that contains the recombinant DNA vector of Claims 1 or 2.
 - 4. An engineered cell line that contains the recombinant DNA expression vector of Claim 1 and expresses Flk-1.
- 5. The engineered cell line of Claim 3 which expresses the Flk-1 on the surface of the cell.
- An engineered cell line that contains the recombinant DNA expression vector of Claim 2 and
 expresses the Flk-1 fusion protein.
 - 7. The engineered cell line of Claim 6 that expresses the Flk-1 fusion protein on the surface of the cell.
 - 8. A method for producing recombinant Flk-1, comprising:
 - (a) culturing a host cell transformed with the recombinant DNA expression vector of Claim 1 and which expresses the Flk-1; and

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- (b) recovering the Flk-1 gene product from the cell culture.
- A method for producing recombinant Flk-1 fusion 5 protein, comprising:
 - culturing a host cell transformed with the (a) recombinant DNA expression vector of Claim 2 and which expresses the Flk-1 fusion protein; and
- (b) recovering the Flk-1 fusion protein from the 10 cell culture.
 - An isolated recombinant Flk-1 receptor protein. 10.
- 11. A fusion protein comprising Flk-1 linked to a 15 heterologous protein or peptide sequence.
- An oligonuclectide which encodes an antisense sequence complementary to a portion of the Flk-1 20 nucleotide sequence, and which inhibits translation of the Flk-1 gene in a cell.
- The oligonucleotide of Claim 12 which is complementary to a nucleotide sequence encoding the amino 25 terminal region of the Flk-1.
 - 14. A monoclonal antibody which immunospecifically binds to an epitope of the Flk-1.
- The monoclonal antibody of Claim 14 which 30 competitively inhibits the binding of VEGF to the F1k-1.
- 16. The monoclonal antibody of Claim 14 which is 35 linked to a cytotoxic agent.

- 17. The monoclonal antibody of Claim 14 which is linked to a radioisotope.
- 18. A method for screening and identifying
 5 antagonists of VEGF, comprising:
 - (a) contacting a cell line that expresses Flk-1 with a test compound in the presence of VEGF; and
 - (b) determining whether the test compound inhibits the binding and cellular effects of VEGF on the cell line,

in which antagonists are identified as those compounds that inhibit both the binding and cellular effects of VEGF on the cell line.

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- 19. A method for screening and identifying agonists of VEGF, comprising:
 - (a) contacting a cell line that expresses the Flk-1 with a test compound in the presence and in the absence of VEGF;
 - (b) determining whether, in the presence of VEGF, the test compound inhibits the binding of VEGF to the cell line; and
- (c) determining whether, in the absence of the VEGF, the test compound mimics the cellular effects of VEGF on the cell line, in which agonists are identified as those test compounds that inhibit the binding but mimic the cellular effects

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20. The method according to Claims 18 or 19 in which the cell line is a genetically engineered cell line.

of VEGF on the cell line.

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- 21. The method according to Claims 18 or 19 in which the cell line endogenously expresses the Flk-1.
- 22. A method for screening and identifying5 antagonists of VEGF comprising:
 - (a) contacting Flk-1 protein with a random peptide library such that Flk-1 will recognize and bind to one or more peptide species within the library;
 - (b) isolating the Flk-1/peptide combination;
 - (c) determining the sequence of the peptide isolated in step c; and
 - (d) determining whether the test compound inhibits the binding and cellular effects of VEGF,

in which antagonists are identified as those peptides that inhibit both the binding and cellular effects of VEGF.

- 20 23. A method for screening and identifying agonists of VEGF comprising:
 - (a) contacting Flk-1 protein with a random peptide library such that Flk-1 will recognize and bind to one or more peptide species within the library;
 - (b) isolating the Flk-1/peptide combination;
 - (c) determining the sequence of the peptide isolated in step c; and
 - (d) determining whether, in the absence of the VRGF, the peptide mimics the cellular effects of VEGF,

in which agonists are identified as those peptides that inhibit the binding but mimic the cellular effects of Flk-1.

- 24. The method according to Claims 22 or 23 in which the Flk-1 protein is genetically engineered.
- 25. A method of modulating the endogenous enzymatic activity of the tyrosine kinase Flk-1 receptor in a mammal comprising administering to the mammal an effective amount of a ligand to the Flk-1 receptor protein to modulate the enzymatic activity.
- 26. The method of Claim 25 in which the ligand to the Flk-1 receptor is VEGF.
 - 27. The method of Claim 25 in which the ligand to the Flk-1 receptor is a VEGF agonist.
 - 28. The method of Claim 25 in which the ligand to the Flk-1 receptor is an antagonist of VEGF.
- 29. The antagonist of Claim 28 that is a monoclonal antibody which immunospecifically binds to an epitope of Flk-1.
 - 30. The antagonist of Claim 28 that is a soluble Flk-1 receptor.
 - 31. The method of Claim 25 in which the enzymatic activity of the receptor protein is increased.
- 32. The method of Claim 25 in which the enzymatic 30 activity of the receptor protein is decreased.
 - 33. The method of Claim 31 in which the ligand stimulates endothelial cell proliferation.

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- 34. The method of Claim 32 in which the ligand inhibits endothelial cell proliferation.
- 35. The method of Claim 32 in which the ligand 5 inhibits angiogenesis.
- 36. A recombinant vector containing a nucleotide sequence that encodes a truncated Flk-1 which has dominant-negative activity which inhibits the cellular effects of VEGF binding.
 - 37. The recombinant vector of claim 36 containing a nucleotide sequence encoding amino acids 1 through 806 of Flk-1.
- 38. The recombinant vector of claim 36 in which the vector is a retrovirus vector.
- 39. The recombinant vector of claim 38 containing
 20 a nucleotide sequence encoding amino acids 1 through 806 of Flk-1.
- 40. An engineered cell line that contains the recombinant DNA vector of Claim 36 and expresses truncated Flk-1.
- 41. An engineered cell line that contains the recombinant vector of Claim 38 or 39 and produces infectious retrovirus particles expressing truncated 30 Flk-1.
 - 42. An isolated recombinant truncated Flk-1 receptor protein which has dominant-negative activity which inhibits the cellular effects of VEGF binding.

43. A method of modulating the cellular effects of VEGF in a mammal comprising administrating to the mammal an effective amount of truncated Flk-1 receptor protein which inhibits the cellular effects of VEGF binding.

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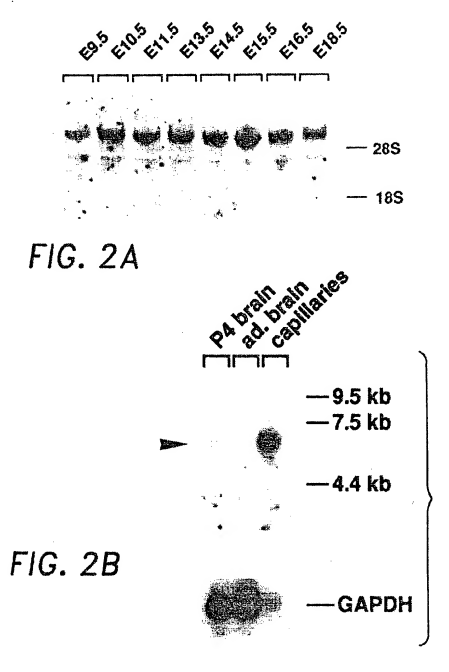
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FIG. 1

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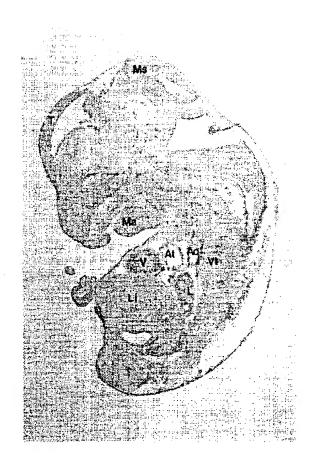


FIG. 3A

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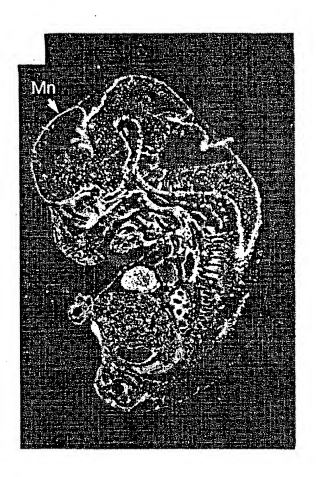


FIG. 3B

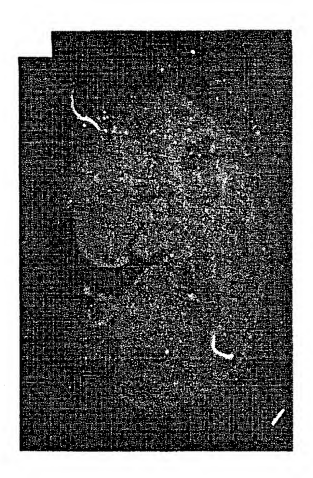


FIG. 3C

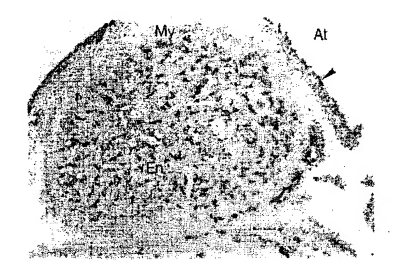


FIG. 4A

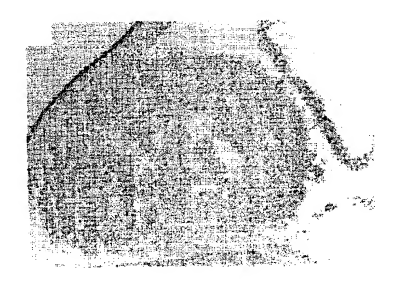
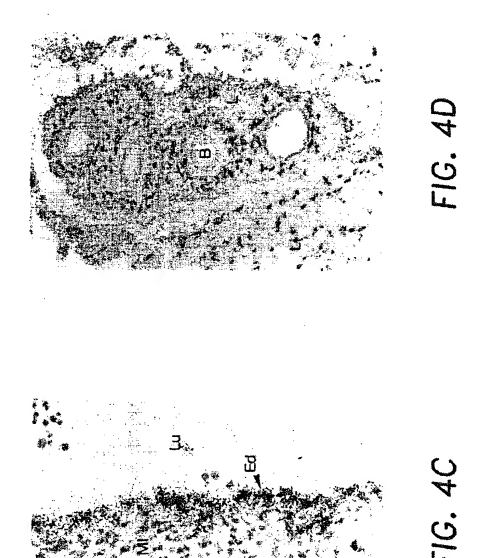


FIG. 4B



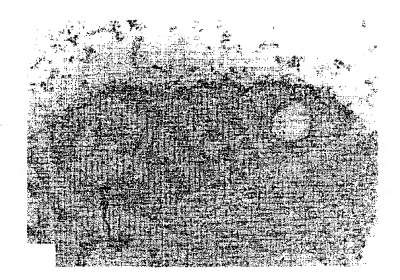


FIG. 4E

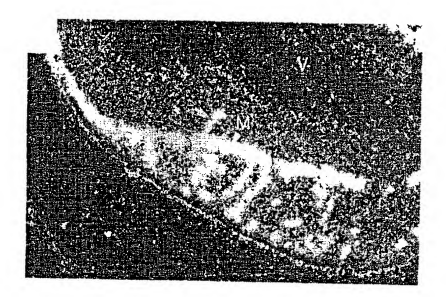


FIG. 5A



FIG. 5B

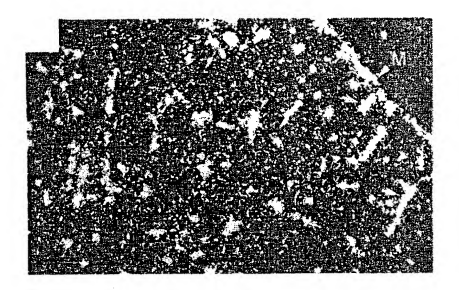


FIG. 5C

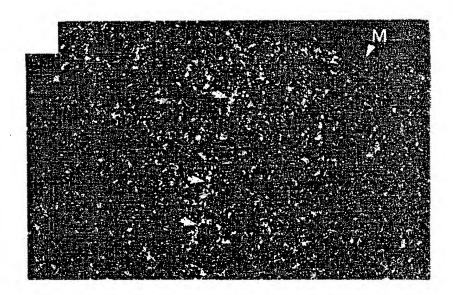


FIG. 5D

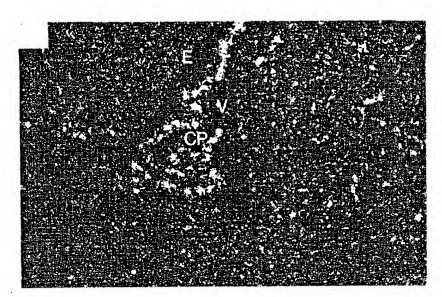


FIG. 6A

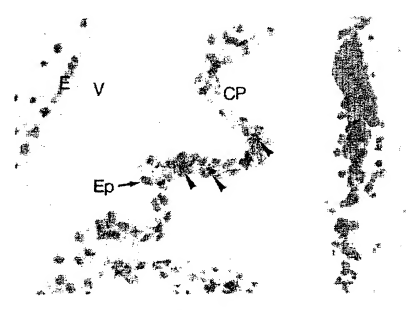


FIG. 6B

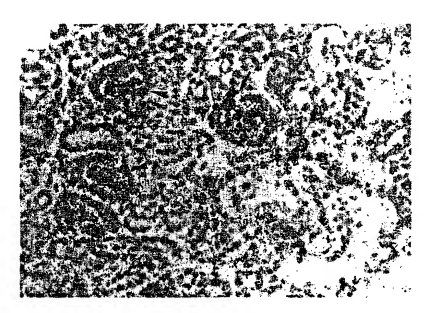


FIG. 7A

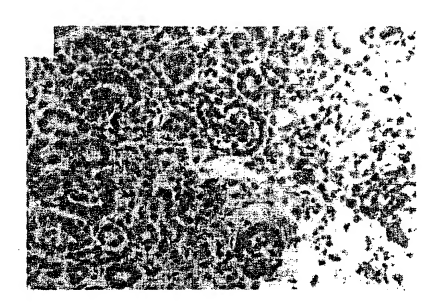


FIG. 7B

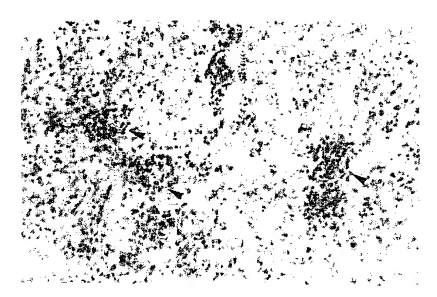


FIG. 7C

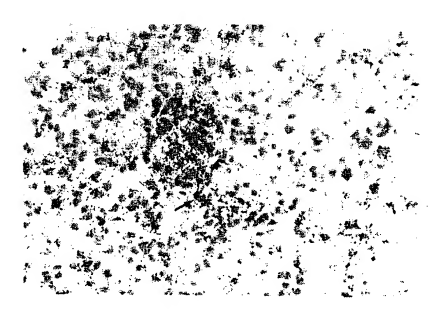
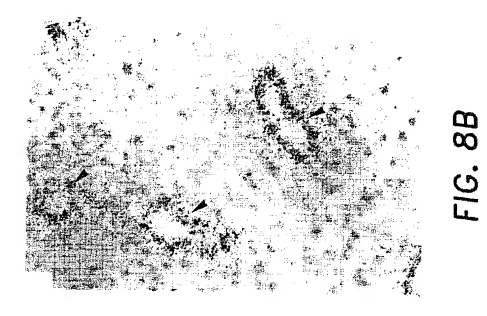
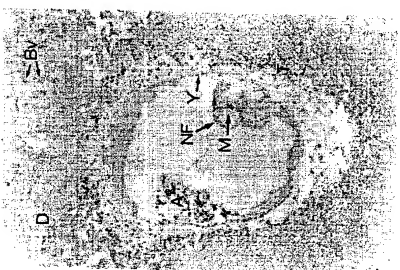
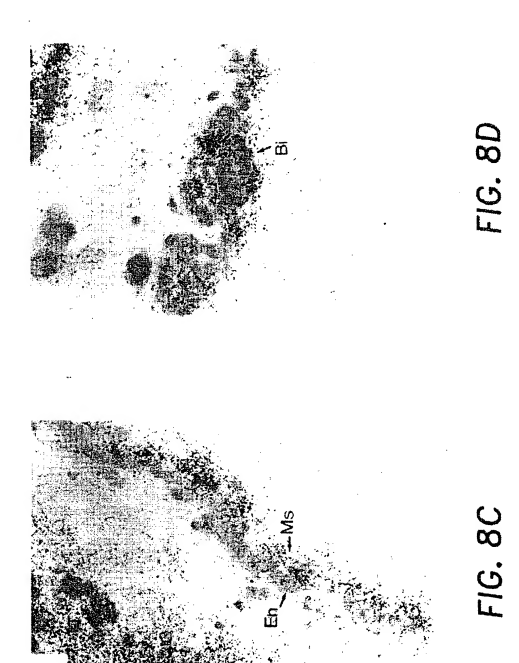


FIG. 7D









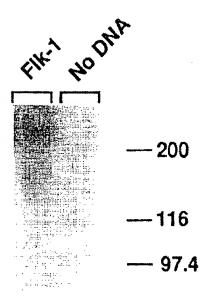
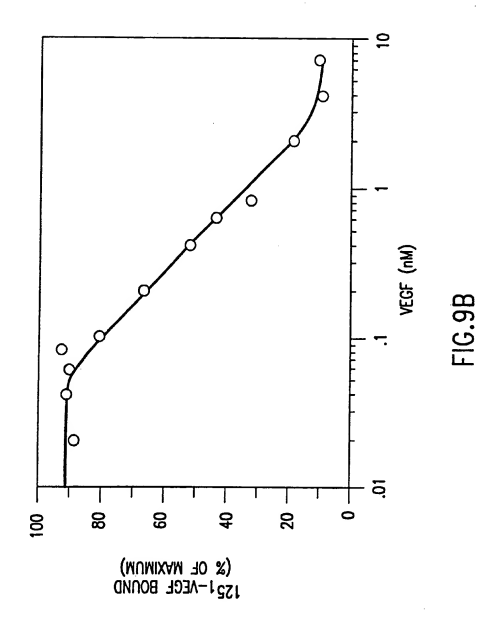


FIG. 9A



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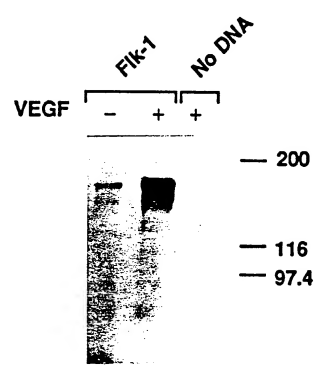


FIG. 10

	1 TATACCCCCAATTCCCCTACCCCCCCCCCCCCCCCCCC	9
9	OCCCATAACCTCCCCCCCCATTCCCCCCCACACCCCCACACCCCCC	16
18	TO CONTROL CON	27
27	M E S K A L L A V A L W F C V H T R A A S V G L T 1 GCCCCCAGGTGCAGGAGCAGCCAGCCCGCTGCTAGCTGTGCCTCTGTGGTTCTGCGTTGGAGCACCCCGAGCCCGCCTCTGTGGGTTTGACT	25 360
26 361	6 G D F L H P P K L S T Q K D I L T I L A N T T L Q I T C R G 1 GGCCATTTICTCCATCCCCCAACCTCAGCACACACACAGACAGACATACTGACAATTTIGGCAAATACAACCCTTCAGATTACTTGCAGCGA	55 450
56 4 51	O R D L D K L M P N A Q R D S E H R V L V T E C G G C D S I CAGCCCCACCTCGACTCGCCTTCGCCCAATGCTCAGCGTGATTCTGAGCAAACCCTATTCGTGACTGAATGCCCCCCGTGCTGACAGTATC	85 540
86 541	FCKTLTTPRVVGNDTGAYKCSYRDVDTAST	115 630
116 - 631	V Y V Y V R D Y R S P F I A S V S D Q H G I V Y I T E N K N GTTTATGTCTATGTTCGGGATTACAGATCACCATTCATCGCCATCGTGGACCAGCATCGCCATCGTGTACATCACCAGAACAAGAAC	145 720
146 721	K T V V 1 P C H G S I S N L N V S L C A R Y P E K R F V P D AAAACTGTGGTGATCCCCTGCCCGACCGTCGATTTCAAACCTCAATGTGTCTCTTTGCCCTACGTATCCAGAAAAGAGATTTGTTCCCGAT	175 810
176 811	G N R I S K D S H I G F T L P S Y M I S Y A G M V F C E A K GGAAACAGAATTICCTGGGACAGGGATAGGCTTTACTCTCCCCAGTTACATGATCAGCTATGCCCGCATGGTCTTCTGTGAGGCAAAG	205 900
206 901	INDKIYQSIMYIYVVVGYRIYDVILSPPHH ATCAATGATGAAACCTATCAGTCTATCATGTACATAGTTGTGGTTGTGGTTGTGGGTTGTAGGATTTATGATG	235 990
236 991	I K L S A G K K L V L N C I A R I E L N V G L D F I M H S P ATTGAGCTATCTGCCGGAGAAAACTTGTCTTAAATTGTACAGCGAGAACAGAGCTCAATGTGCGGCTTGATTTCACCTGCCACTCTCCA	265 1080
266 1081	PSKSHHKKIVNRDVKPFPGTVAKMFLSTLTCCTTCCAAAGTCTCATAAGAAGATTGTAAACCCCGATGTGAAACCCTTTCCTCCGGACTGTGGGGGAAGATGTTTTTGAGCACCTTTGACA	295 1170
296 1171	I E S V I K S D Q G E Y I C V A S S G R N I K R N R I F V R ATAGAAAGIGTGACCAAGAGGACAAGAGGAATAGAACATTIGICOGA	325 1260

FIG.11A

326	¥	ł	1	Ī	K	P	F	'		A	F	G	S	G	V	K		5 1	L '	V	E	A	1	٧	G	S	Q	٧	R	1	F	, 1	/ K	35
1261	G	110	X	AC	₩	GC	CIT	11/	۱TI	GC	III	Œ	STAC	CTG	CGA	TGA	AA]	CT	TG	GTG	GAA	100	CAC	AG 1	CC	CCA	GTC	AAG	ΙΩ	GAA	ICC	CTO	TGAA	3 135
356 1351	Y	L	TC	s ag	Y Tta	P CCC	À Cag	CTC	CI	D Gai	I Tat	K	N VATO	Y ST	R ACA	N Gaa	ATG	GA/	1000 1000	P CCC	I ATT	E Ga	S CTC	N Cav	Y CT/	T NCAI	M AK	I Tga	V TTG	G TTG	CCC	ATC	(L Vaact(38 144
386 1441	T A(i Ca	TC	M at(K CCA	V AG1	T [GA	K CTG	A A	r Ag/	D VCA	A TGC	Q	N AA	Y ACT	T ACAI	V	TCA	l ITC(L CTC	T ACC	N 'AA(P	i Cat	S TT(N CAA	E IGG	K NGA	Q NC	S Aga	H	M ACA	I V Togto	41:
416 1531	S	L CTC	TC	V GT1	V IGT	K Gav	V ITG	P TCC	CA	P CCC	Q Xa	I Cat	C	E TC/	K NGA	a Vag(l XI	1 Tga	itci	s icc	P CCT	M ato	D Xa	S TIC	Y CT/	Q VCC/	Y GT/	G ATO	T Ga(M CA	Q TGC	Y Aga	L Cattg	44! 162
446 1621	T AC	C AT	CC	r IC/	V GT	Y Cta	A CC(N XX	AC	P XI	P CC	L CCI	H GCA	H 00/	I VCA1	0	N GT	Y CCT	AC1	l [GG	Q (L CT <i>A</i>	E VCA	E NGA	A AGC	C CTG	S CT(Y XTA	R CAC	P AC	0 2003	Q 200	T AMACA	47: 171(
476 1711	S AG	P CC	r Tx	r Tat	A GC	C TTG	K Taj	E WG	ļ AA	(TGC	R Vaga	H Aca	V CCT	E CCA	D GG/	F ITTI	Q 221	G ACC	000	GAV	WC)	K Aac	I Ato	E Xia	V Agt	T Cac	K Ca4	N AA4	Q XX	Y ATA	A ATG	m L	I Igatt	505 1800
506 1801	K Ga	G ACX	N W	(IAA	n aa(K Caa	T AAC	V TG	ia/	S NGT.	T ACC	L CCT(V CCT	I Cat	Q CC/	A AGC	A TCI	N CCA	V ACG	TG	CAC	A GCC	L 11(Y Ta	K Caa	C atg	E Tga	A GCC	I ATC	N XAA	K Aa	A CO	G XCCA	535 1890
536 1891	R CC.	G ACX	E Sag	AG	R AGO	V GT	I Cat	S CT(F Cl	: 110	H Ca1	V IGT(I Gat(R CAG	C CCC	P TCC	E TG/	I WA	T ATI	CTO	/ () `AA	P CC1	A CC	A ICO	Q CCA	P GCC	T AAC	E TGA	Q CC/	E GG/	S (CA)	V STGTG	565 1980
566 1981	S	L XI	L	īG	C TCC	T AC	A TCC	D AG/	R NCA	R (N AA1	T ACC	F STT	E Iga	N Gaa	L CCT	T CAC	N CT(Y CI.	K ACA	AGC		G CCC	S :TC/	Q VCA	A CCC	T AAC	S ato	V GCT	H CC/	N CAT	G ()()	E XCAA	595 2070
596 2071	S TC/	L VCI	T CA	CA	P CCA	V Gt	C ITG	K Ca/	N IGA	I I AC'	L TTG	D Gai	A IGC1	L	N TTG	K Gaa	L AC1	M Ga/	G VIG	T GCA	CCA	l I	F TTT	S TC1	N Aad	S CAG	T	N AAA	D Tga	I Cat	L CT1	I Gat	V TGTG	625 2160
626 2161	A GC/	F \TT	Q TC	I AG/	N NAT	A CCI	S CTC	L ICI	Q CC	ACC) CAC	Q Caf	G GGC	D Gai	Y CTA	V TGT	C TTG	S	A CTG	Q CTC	AAG) AT/	K AAG	K AAC	T ACX	K Xaa	K Saa	r Ng	H Aca	C TTG	L CCT	V GGT	K Caaa	655 2250
656 2251	Q CAO	i CT	I Ca	TC/	I ATC	L CT/	K Igai	R CCC	M CA	TGC	i Xa	P CCC	H ATG	I AT(T CAC	G XX	N Aaa	L ICT	2 (30)	N Aga	Q atc	AG	T ACA	T ACA	T ACC	I ATI	Q CCC	E Ca	T Gaci	I Cat	H TGA	V Agt	T Gact	685 2340
686 2341	C TGC	P CC	A ACI	CAI	S ICT	C GG/	N WA	P TCC	TA	F CCC	CA	N Cac	I ATT	T ACA	K VTO:	F STT(K Caa	D Aga	N Cav	3 204	T Aga	j ca	CIO	V GTA	E Gaa	D GA1	S TC/	C C	I ATI	V IGT	L ACT	R Gag	D AGAT	715 2430
716 2431	G CCC	N Aa	R	N GA	VQ	L CTG	T AC1	I Tat	R CCI	R GCA	CCI	V GTG	R ACG	K Aac	E Ca(D Gai	G IGG	G ACC	L CC1	Y ICT/	T ACAI	(133	; (Q CAO	A GCC	C ICC	N 'AA1	V GT(L XII	G GC	C CTG	A TGC	r aaga	745 2520

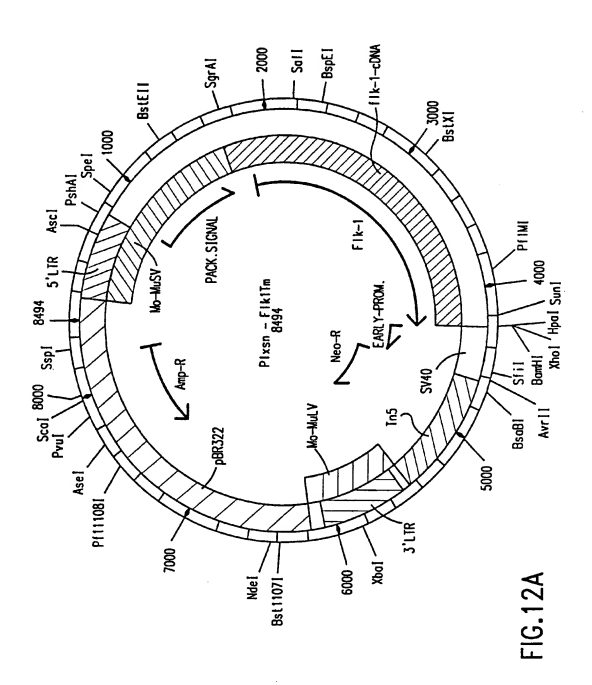
FIG.11B

	A E T L F I I E G A Q H K I N L E V I I L V G I A V I A M F GOOGAGACCTCTTCATAATAGAAGGIGCCCAGGAAAAGACCAACTTGGAAGTCATTATCCTCCTCCCCACTGCAGTGATTGCCCATGTTC	775 2610
776 2 611	F M L L L V I V L R T V K R A N H G K L K T G Y L S I V M D TICTOGCTCCTTCTTGTCATGTCCTACCGACCGTTAAGCCCCCAATGAAGCCCGAACTGAAGACAGCCTACTTGTCTATTGTCATGGAT	805 2700
	F D K L P L D H R C K E L P Y D A S K N E F P R D R L K L G CCAGATGAATTGCCCTTGCGTGGAACCTGCACTTGCCTTATGATGCCACCAAGTGCGAATTCCCCACCGACCCCCTGAAACTAGGA	835 2790
	K F L G R G A F G Q V I E A D A F G I D K T A T C K T V A V AAACCTCTTGCCCGCGGTGCCTTCGGCCAAGTGATTGAGGCAGACGCTTTTGGAATTGACAAGACAGCAACACGCAAAACAGTAGCCGTC	865 2880
	K M L K E G A T H S E H R A L M S K L K I L I H I G H H L M AAGATGTTGAAAGGAGGAGCAACACACACGCGAGCATCCACCCTCATGTCTGAACTCAAGATCCTCACCATTGGTCACCATCTCAAT	895 2970
	${\tt V}$ ${\tt V}$ ${\tt N}$ ${\tt L}$ ${\tt G}$ ${\tt A}$ ${\tt C}$ ${\tt I}$ ${\tt K}$ ${\tt P}$ ${\tt G}$ ${\tt G}$ ${\tt P}$ ${\tt L}$ ${\tt N}$ ${\tt V}$ ${\tt I}$ ${\tt V}$ ${\tt E}$ ${\tt F}$ ${\tt C}$ ${\tt K}$ ${\tt F}$ ${\tt G}$ ${\tt N}$ ${\tt L}$ ${\tt S}$ ${\tt I}$ ${\tt Y}$ ${\tt G}$ ${\tt G$	925 3060
926 3061	L E G K R N E F V P Y K S K G A R F R Q G K D Y V G K L S V TTACCCCCCAGGCCAAGGACATGAATTTGTTCCCTATAAGACCAAAGGCCCAGGCTTCCCCCAGGCCAAGGACTACGTTGCGGACCTCTCCGTG	955 3150
	D L K R R L D S I I S S Q S S A S S G F V K H K S L S D V E CAICTGAAAAGACGCTTCGACACCATCACCAGCAGACCTCTCCCAGCTCTCCCCCTCACCCTTTGTTGAGCAGAAATCCCTCAGTGATGTAGAG	985 3240
	K K K A S K K L Y K D F L T L K H L I C Y S F Q V A K G M E GAAGAAGAAGCTTCTGAACAACTGTACAAGCACTTCCTGACCTTGCAGCATCTCATCTGTTACAACTTCCAAGTGCCTAAGCCCATGGAG	1015 3330
	F L A S R K C I H R D L A A R N I L L S E K N V V K I C D F TICTICCCATCAACGAACGTGTATCCCACCACCACCACCACCACCACCACCACCACCACCA	1045 3420
-	G L A R D I Y K D P D Y V R K G D A R L P L K K M A P E T I GCCTTGGCCCCGGACATTTATAAAGACCCCGATTATGTCAGAAAAGGACATGCCCCGGACTCCCTTTGAAGTGGATGCCCCCCGAAACCATT	1075 3510
3511	F D R V Y I I Q S D V N S F G V L L N E I F S L G A S P Y P THIGACAGAGTATACACAATICAGAGCGATGTGTGCTCTTTCCGTGTGTTGCTCTCCCGAAATATTTTCCTTACGTCCCTCCC	1105 3600
	G V K I D E E F C R R L K E G T R M R A P D Y T T P E M Y Q	1135

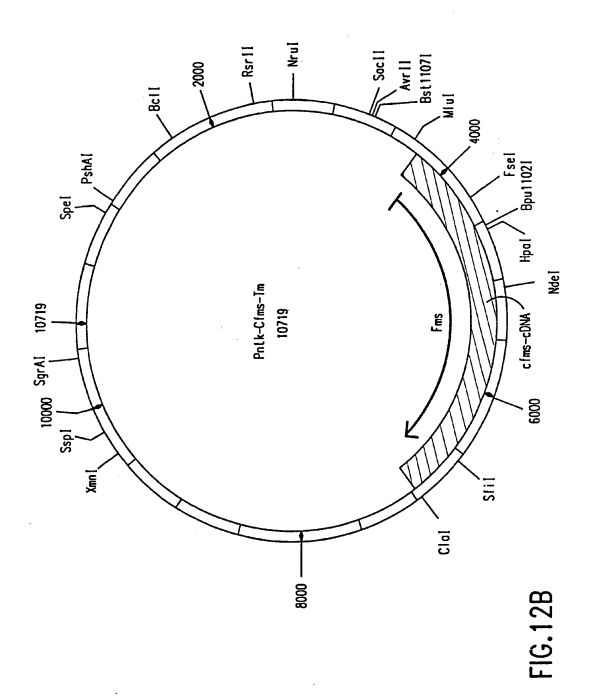
FIG.11C

3691	ACCATOCTOGACTOCTOCCATGAGGACCCCAACCACAGACCCTCGTTTTCAGAGTTGGTTG	3780
	A Q Q D G K D Y I V L P M S E T L S M K E D S G L S L P T S GCCCAGCAGGAGAGGATTCTGCACTATCTTCCAATGTCAGAGACACTGAGCATGGAAGACGATTCTGCACTCTCCCTGCCTACCTCA	1195 3870
	P V S C M E E E H V C D P K Y H Y D N T A G 1 S H Y L Q N S CCTG+TTCCTGTATCGACGAGGGAGGGAGGGAGGGACCCCAAATTCCATTATGACAACACGCACG	1225 3960
	K R K S R P V S V K T F H D I P L E E P E V K V I P D D S Q AAGCGAAAGAAGTGAGGCCCAGTGAGTGACACCAGATGACACCAGATGACACCAGA	1255 4050
	T D S G M V L A S E E L K T L E D R N K L S P S F G G M M P ACAGACAGTGGGATGGTCCTTGCATCATCAGAAGAGCTGAAAACTCTGGAAGACACAAATTATCTCCATCTTTTGGTGGAATGATGCCC	1285 4140
	S J S R E S V A S E G S B Q T S G T Q S G T G S D D T D T T AGTAAAAGCAGGGGGTGTGGGCTGGGAAGCCTCCAACCAGACCACCTGCCTACCAGTCTGGGTATCACTCAGATGACACAGACACCACC	1315 4230
	Y Y S S D E A G L L K M Y D A A Y H A D S G T T L Q L T S C GIGIACTCCACGCACCACGCACCACCACCTCCACCTCCTCT	1345 4320
	L N G S G P V P A P P P T P G N H E R G A A + TTANATGGAAGTGGTCCTGTCCCCGCCCCCCCCCCCCCCC	1367 4410
4411	CACCACCOGGAAGTAGCCACATTTGATTTTCATTTTTGGAGGAGGGACCTCAGACTGCAAGCAGCTTGTCCTCAGGGCATTTCCAGAGAA	4500
4501	GATGCCCCATGACCCAAGAATGTGTTGACTCTACTCTCTTTTCCATTCATT	4590
4591	CAGTTAAAGCAAAAGACTTTCAAACAOGTGGACTCTGTOCTOCAAGAAGTGGCAACOGCAACCTCTGTGAAACTGGATCGAATGGGCAATG	4680
4681	CTTTGTGTGTGAGGATGGGTGAGATGTCCCAGGCCCCAGTCTGTCT	4770
4771	TAACTGTCGCATGTCCACTGGCAGCAAGGAAGGCCCAAGTCGCTCGGACAGCCGTTGCAGCCTGCACATCCATTGTGCTCGCTC	4860
4861	AGGTGGGCTTGTGGCCTGTCAGGAAACCCAAACGCCCCCCCC	4950
4951	COCAGTICCCTGTCCCCTTTCCTACTCCTAATCAGAGTTCCTTCCCCGCACTCTTACCTGTCTCCTCCCCCCCACCAAACCAAATGATC	5040
5041	CAGCTTGCTCCTTCCTCATCTCTCAGGCTGTGCCTTAATTCAGAACACCAAAAGAGAGGAACGTCCCCAGAGCCTCCTGACCGCCCGAA	5130
5131	CAATTGTGACAACAGAACAGAAACTCACCGTTTCTGCTGCGTCGACACCCACGTGCCCCCCTCCTGCCACGTCTGACCGTTCTCTGTCAA	5220
5221	GTGGCCCTAAACGCTCAGCCTGGTGTTCTTCCTCTATCTCCACTCCTGTCAGCCCCCCAAGTCCTCAGTATTTTACCTTTTGTGGCTTCCT	5310
5311	CATCCCAGAAAAATCTTAATTCGTTGCTTTCCTCTCCAGATAATCACTACCCAGATTTCGAAATTACTTTTTTACCCGACGTTATGATAAC	5400
5401	ATCTACTGTATCCTTTAGAATTTTAACCTATAAAACTATGTCTACTGGTTTCTGCCTGTGTGCTTATGTT	5470

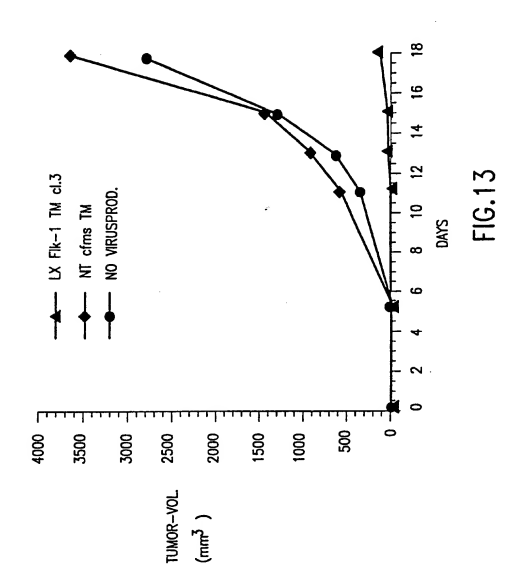
FIG.11D SUBSTITUTE SHEET



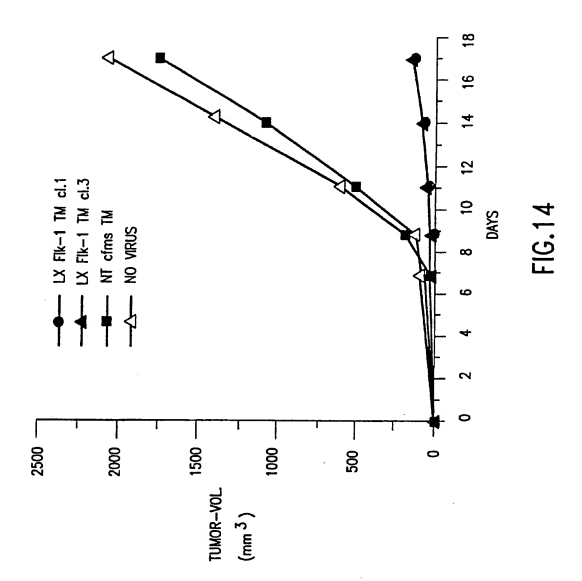
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utional Application No PCT/EP 93/03191

A. CLASSIFICATION OF SUBJECT MATTER
IPC 5 C12N15/12 C07K13/00 C12P21/08 C12N15/86 C12Q1/68 G01N33/567 A61K37/02 C12N15/62 According to International Patent Classification (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classification system followed by classification symbols) IPC 5 CO7K C12N A61K Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted thring the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Relevant to claim No. Citation of document, with indication, where appropriate, of the relevant passages Category ' 1-11 PROCEEDINGS OF THE NATIONAL ACADEMY OF X SCIENCES OF USA. vol. 88 , October 1991 , WASHINGTON US pages 9026 - 9030 MATTHEWS, W. ET AL.; 'A receptor tyrosine kinase cDNA isolated from a population of enriched primitive hematopoietic cells and exhibiting close genetic linkage to c-kit. see the whole document 1-11. WO,A,92 17486 (TRUSTEES OF PRINCETON 14-37 UNIVERSITY, US) 15 October 1992 12,13, Y see the whole document 38,39,41 -/--Further documents are listed in the continuation of box C. Patent family members are listed in annex. X Special categories of cited documents: "I" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the "A" document defining the general state of the art which is not considered to be of particular relevance invention P. earlier document but published on or after the international "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone filing date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such docu-ments, such combination being obvious to a person skilled "O" document referring to an oral disclosure, use, schibition or document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report 115 -04- 1994 11 March 1994 Name and mailing address of the ISA **Authorized** officer Buropean Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk Tel. (+ 31-70) 340-2040, Tx. 31 651 epo nl, Faz: (+ 31-70) 340-3016

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C.(Continue Category	ation) DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
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Y	WO,A,92 03459 (SLOAN KETTERING INSTITUTE OF CANCER, US) 5 March 1992 see the whole document	12,13
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P,X	CELL vol. 72 , 26 March 1993 , CAMBRIDGE, NA US pages 835 - 846 MILLAUER, B., WIZIGMANN-VOOS, S., SCHNURCH, H., MARTINEZ, R., MOLLER, N.P., RISAU, W., AND ULLRICH, A.; 'High affinity VEGF binding and developmental expression suggest Flk-1 as a major regulator of vasculogenesis and angiogenesis.' see the whole document	1-43
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PCT/EP 93/03191

Box 1 Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This international search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. X claims Nos.: hecause they relate to subject matter not required to be searched by this Authority, namely: Remark: Although claims 25-28, 31-35, 43 are directed to a method of treatment of the human/animal body as well as diagnostic methods (Rule 39.1 (iv) PCT) the search has been carried out and based on the alleged effects of the compound/composition.
2. Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:
3. Claims Nos hecause they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
The International Searching Authority found multiple inventions in this international application, as follows:
As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.
2. As all searchable claims could be searches without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:
4. No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark on Protest The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.

Information on patent family members

F sational Application No PCT/EP 93/03191

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